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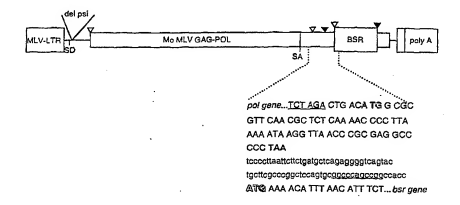
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(54) Title: EXPRESSION SYSTEMS



Schematic structure of CeB expression vector

(57) Abstract

The invention relates to new expression systems and in particular to an expression system in which a gene of interest is expressed at an optimal level. The invention provides a recombinant expression vector comprising a gene of interest and a selectable marker gene, wherein the selectable marker gene is arranged downstream of the gene of interest and a stop codon associated with the gene of interest is spaced from a start codon of said selectable marker gene at a distance which is sufficient to ensure that translation reinitiation is required before said selectable marker protein is expressed from the corresponding MRNA. Examples of such expression systems are vector viral packaging cell lines and a number of preferred cell lines have been identified.

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Expression systems

The present invention relates to new expressions systems, and in particular to expression systems in which a gene of interest is expressed at an optimal level. Particular examples of such expression systems are retroviral packaging cell lines and a number of preferred cell lines have been identified.

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The ability of eukaryotic and prokaryotic ribosomes to reinitiate translation at an internal start codon within an mRNA sequence has previously been recognised. Studies have been reported in which the efficiency of the process, which is generally regarded as being low, has been connected with the length of the intercistronic sequence (Kozak (1987) Mol. Cell Biol. 7, 3438-3445). Selection of this sequence or spacer as 70bp in length, and containing no other start codons, has been previously reported as being optimal for reinitiation in a eukaryotic cell line (Cosset F-L., Virology (1991) 185, 862).

The applicants have found a way in which the inefficiency associated with the translation reinitiation process can be used to good effect.

According to the present invention there is provided a recombinant expression vector comprising a gene of interest and a selectable marker gene, wherein the selectable marker gene is arranged downstream of the gene of interest and a stop codon associated with the gene of interest is spaced from a start codon of said selectable marker gene at a distance which is sufficient to ensure that translation reinitiation is required before said selectable marker protein is expressed from the corresponding mRNA.

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The invention further provides a process for producing cell lines in which a gene of interest is expressed, which process comprises transforming host cells with an expression vector comprising said gene of interest and a selectable marker gene, wherein the selectable marker gene is arranged downstream of the gene of interest and a stop codon associated with the gene of interest is spaced from a start codon of said selectable marker gene at a distance which is sufficient to ensure that translation re-initiation is required before said selectable marker protein is expressed from the corresponding mRNA, and selecting those cells where expression of the selectable marker gene may be detected.

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Since re-initiation of translation is a relatively
inefficient process, this means that the selectable marker
protein will be expressed at lower levels than the product
of the gene of interest. When the marker protein is
expressed at detectable levels, the gene of interest will be
expressed at higher levels. This will ensure that during
the subsequent selection procedure, only those cell clones
which express the gene of interest at higher or optimal
levels will survive. Low expressing clones will be
eliminated by the selection process.

Cells transformed with the above-described expression vectors form a further aspect of the invention.

The host cells are suitably eukaryotic or prokaryotic host cells, preferably eukaryotic host cells.

The number of nucleotides in the space between the stop codon of the gene of interest and the start codon of the selectable marker will suitably be in the range of from 20-200 nucleotides, preferably from 60-80 nucleotides, even more preferably 70-80 nucleotides.

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The vectors used in the process of the invention may be any of the known types, for example expression plasmids or viral vectors.

Selected cells may be cultured and if required, the protein product of the gene of interest isolated from the culture using conventional techniques. Alternatively, expression of the gene of interest may result in other desired effects, for example, where the gene of interest is included as part of a viral packaging construct.

Some experimental and clinical gene transfer protocols require the design of gene transfer vectors suitable for in vivo gene delivery (Miller, A.D. 1992. Nature 357:455-460). Retroviral vectors are attractive candidates for such applications, because they can provide stable gene transfer and expression (Samarut J. et al., Meth. Enzymol. in press) and because packaging cells have been designed which produce non-replication competent viruses (Miller A.D (1990) Hum Gene Ther. 1 5-14). However currently available recombinant retroviruses suffer from a number of drawbacks.

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Packaging cell lines provide <u>in trans</u> the retroviral proteins encoded by the <u>gag</u>, <u>pol</u>, and <u>env</u> genes required to obtain infectious retroviral particles. The <u>gag</u> and <u>pol</u> products are respectively the structural components of the virion cores and the replication machinery (enzymes) of the retroviral particles whereas the <u>env</u> products are envelope proteins responsible for the host-range of the virions and for the initiation of infection and for sensitivity to humoral factors. An ideal packaging cell line should produce retroviruses that only contain the retroviral vector genome, and absolutely no replication-competent genomes or defective genomes encoding some of the viral structural genes.

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A number of packaging cell lines designed for human gene transfer have been designed in the past by introducing plasmid DNAs which contain "helper genomes" encoding gag, pol and/or env genes into cells.

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Retroviral packaging cell lines are cells that have been engineered to provide in trans all the functions required to express infectious retroviral vectors. A helper genome (or construct or unit), is herein also referred to as "retroviral packaging construct (or unit)" or "packaging-deficient construct (or genome unit)" or "gag-pol/env expression plasmids".

Much efforts has been made to design strategies to optimize
the helper-genomes in order (i) to get the highest
production of retroviral packaging functions (which
correlates which infection titers of retroviral particles)
and (ii) to minimise the chance that the helper genome can
be transmitted via the viral particles (which may lead to
emergence of unwanted retroviral forms).

The first of these packaging cell lines used full length retroviral genomes as helper genomes that had been crippled for important cis-regulated replicative functions (reviewed in Miller, Hum. Gene. Ther. 1:5-14 1990). In order to reduce the possibility of occurrence of replication-competent viruses and of transfer of virus structural genes, a second generation of safer packaging cell lines has been designed by using two separate and complementary helper genomes which express either gag-pol or env and are packaging-deficient (Miller supra).

The cells into which these helper genomes were introduced were isolated by cotransfecting them with plasmids encoding selectable markers. However, as no selection was applied on

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the packaging-deficient retroviral genome itself, the helper functions can be lost during the passages of the cells in culture and the current packaging systems provide limited titers of infectious retroviral vectors, usually only of the order of 10^5-10^6 infectious units i.u/ml. Indeed the cotransfection with a plasmid encoding a selectable marker does not directly select the best gag-pol-env-expressing cells.

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10 The invention further provides a retroviral packaging cell line comprising a host cell transformed with (i) a packaging deficient construct which expresses a viral gag-pol gene and a first selectable marker gene, and/or (ii) a packaging-deficient construct which expresses a viral env gene and a second selectable marker gene; wherein a start codon of the first and second selectable markers are spaced from the stop codons of the viral gag-pol gene and the viral env gene respectively by a distance which ensures that reinitiation of mRNA translation is required for expression of marker protein product of said first and/or second selectable marker gene.

The retroviral packaging cell line may be obtained by the above described process which will involve selecting transfected cells which express said first and/or second marker genes.

By using helper constructs which are directly selectable and which provide for high expression of the viral gene, high titre retroviral vectors may be obtained.

Helper constructs for use in the process form a further aspect of the invention.

35 The retroviral vectors prepared from the conventional

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packaging cell lines are usually not contaminated by replication-competent retroviruses (RCRs). However, recombinant amphotropic murine retroviruses have been shown to arise spontaneously from certain packaging cell lines. The generation of such RCRs involves recombination at least between gag-pol/env packaging sequence and vector sequences (Cosset et al., Virology, (1993) 193:385-395).

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Recombinant RCRs have been associated with the development of lymphomas in some severely immunosuppressed monkeys 10 (Donahue et al., J. Exp Med (1992) 176: 1125-1135). In addition, retroviral vector preparations may also contain, at low frequencies, retroviruses coding for functional envelope glycoproteins (Kozak and Kabat, 1990, J. Virol. 64: 3500-3508) or for gag-pol proteins. Although the 15 pathogenicity of these gag-pol or env recombinant retroviruses is probably low, more evolved recombinant retroviruses with higher pathogenic potential may occur when injected in vivo, by recombination and/or complementation of the initial recombinant viruses with some endogenous 20 retroviruses.

In a preferred embodiment of the retroviral packaging cell lines of the invention, the overlapping sequences between the genomes of the retroviral vector and the helper construct are reduced, for example as compared to constructs such as CRIPenv and CRIPAMgag (Danos et al., Proc. Natl. Acad. Sci USA 85: 6460-6464). In particular, the viral sequences in the helper construct are reduced, for example, not only the packaging sequence but also the 3' Long Terminal Repeat(LTR), the 3' non-coding sequence and/or the 5'LTR may be eliminated.

The possibility of generation of such RCRs and recombinant retroviruses can be reduced by reducing the overlapping

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sequences between the genomes of both the retroviral vector and the helper construct.

Conventional retroviral vectors are strongly inactivated by human serum which makes them of limited or no use for in5 situ gene transfer in gene therapy applications. previously been shown that inactivation by complement in human serum is controlled by the cell line used to produce the virions and by viral envelope determinants (Takeuchi et al., J. Virol (1994) 68:8001-8007). In particular, 10 inactivation is caused by some properties of the cell lines that have been used to construct the packaging cells (NIH-3T3) and also by viral determinants located in the retroviral envelope as shown (Takeuchi et al., J. Virol (1994) **68:**8001-8007). <u>In vivo gene delivery is an important</u> 15 goal for a number of human gene therapy strategies.

The applicants have found that certain cell lines form preferred packaging cell lines.

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Particularly preferred packaging cell lines are the HT1080 line, the TE671 line, the 3T3 line, the 293 line and the Mv-1-Lu line. One example of retroviral packaging cells that will produce complement-resistant virus comprise human HT1080 cells and express RD114 envelope. Such cells form a preferred aspect of the invention.

Packaging cell lines according to the invention provide 50-100 fold increased titers of retroviral vectors as compared to conventional packaging cell lines. Retroviral vectors provided by these new cells are safe, in terms of generation of RCRs, and considerably more resistant to inactivation by human complement.

Packaging cell lines according to the invention may be able

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to transduce helper-free, human complement-resistant retroviral vectors at titers consistently higher than 10^7 i.u./ml.

Suitable semi-packaging cell lines in accordance with the invention are those which express only the gag-pol genes. Such cell lines may suitably be derived from TE671, MINK Mv-1-Lu, HT1080, 293 or NIH-3T3 cells by introduction of plasmid CeB (the MoMLV gag-pol expression unit).

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Particularly preferred expression vectors in accordance with the invention for use in retroviral packaging cell lines are those which include MLV gag and pol genes such as CeB. Other plasmids may include gag and pol genes from other retroviruses or chimeric or mutated gag and pol genes.

Various viral and retroviral envelope genes may be included in the plasmids such as MLV-A envelope, GALV envelope, VSV-G protein, BaEV envelope, RD114 envelope and chimeric or mutated envelopes. Plasmids which include the RD114 env gene such as FBdelPRDSAF as illustrated hereinafter, provide one example of suitable constructs.

The novel retroviral packaging cells described hereinafter,

have been designated FLY cells, and may be designed for in

vivo gene delivery.

Considerable variations were found between the various cell lines screened for their ability to release type C mammalian retroviruses. In addition, few cell lines were able to produce retroviruses completely resistant to human complement. Based on these two criteria, human fibrosarcoma HT1080 and rhabdomyosarcoma TE671 cells were selected for optimum construction of packaging cells.

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Other studies have shown the importance of endogenous retrovirus expression in the generation of recombinant retroviruses from retroviral packaging lines (Ronfort et al., Virology, (1995), 207, 271-275, Vanin, E.F.et al., J Virol (1994) 68:4241-4250.). The co-packaging of an 5 endogenous genome and a vector can lead to emergence of recombinant retroviruses (Vanin et al., supra). Recombination involves template switching during reverse transcription of such hybrid retroviruses (Hu et al., Science, (1990) 250:1227) and homologies between the two genomes considerably enhance the frequency of reverse transcriptase jumps (Zhang et al., J. Virol. (1994) 68: 2409-2414). Therefore an ideal packaging cell line should not express endogenous MLV-like (or type C retrovirus-like) retroviral genomes which can be packaged by type C gag proteins (Scadden et al., J. Virol. (1990) 64: 424-427, Torrent et al., J. Mol. Biol. (1994) 240 434-444).

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Packaging of human endogenous retroviral RNA was not detected in TELCeB and FLY packaging cells when virion 20 associated RNA was analysed by RT-PCR using generic primers. HT1080- and TE671 derived packaging cell lines may be safer in this respect than those generated from NIH3T3 cells, such as GP+EAM12 cells, which are known to express and package sequences related to type C retroviruses (Scadden et al. 25 supra).

To generate the FLY packaging cell lines, HT1080 cells were transfected with gag-pol and env expression plasmids designed to optimise viral protein expression. selection for viral gene expression was achieved in accordance with the invention by expression of a selectable marker gene by re-initiation of translation of the mRNA expressing the viral proteins. This strategy resulted in packaging cell lines capable of producing extremely high

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titer viruses. Furthermore, long-term expression of packaging functions can be maintained in these cells. Many unnecessary viral sequences were eliminated from the packaging constructs to reduce the risk of helper virus generation; indeed the final packaging cells did not produce helper virus, in that no replication competent virus (RCR) could be detected per 107 vector particles.

The FLY packaging cells described herein are safer than, for example, psiCRIP cells, at least for generation of env 10 recombinant retroviruses as is illustrated in Table 4 hereinafter, probably because less retroviral sequences overlapping with the vector were present in the present envexpression plasmid. Few reports have addressed the question of the characterization of recombinant retroviruses (RVs) 15 (Cosset, F.L., et al., Virology (1993) 193:385-395). It is possible that such RVs could not be detected in previous packaging cell lines due to lower overall titers. RVs are defective in normal cell culture conditions but are likely 20 to evolve to replication competent viruses if they are allowed to replicate in cells complementing their expression like co-cultivated packaging cells (Bestwick et al., Proc. Natl Acad Sci USA, (1988) 85: 5404-5408, Cosset et al., (1993) supra).

In preferred retroviral packaging systems according to the invention, RVs are eradicated for example by removal of viral LTRs from the packaging construct.

Consistent with our previous studies (Takeuchi, Y., et al., J Virol (1994) 68:8001-8007), LacZ(RD114) and lacZ(MLV-A) pseudotypes produced from HT1080 and TE671cells were more resistant to human complement than LacZ(RD114) or LacZ(MLV-A) pseudotypes produced by 3T3 of dog cells. It was therefore decided to use RD114 and MLV-A env genes to

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generate recombinant virions with MoMLV cores.

The sequence of RD114 env gene was determined and is shown in Figure 4. It was found to be very close to BaEV (baboon endogenous virus) a type C retrovirus (Benveniste, R.E.et 5 al., Proc. Natl. Acad. Sci. USA (1973) 70:3316-3320; Kato, S.et al., Japan. J. Genet. (1987) 62:127-137) with an envelope gene displaying similarities to the external part of type D simian retroviruses (SRVs). RD114 uses the SRV receptor on human cells (Sommerfelt & Weiss, Virology 10 (1990) 176:58-69; Sommerfelt, M.A. et al., J Virol (1990) 64:6214-6220) making the FLY packaging cells with RD114 envelope capable of generating virions with different tropism. Retroviral vectors prepared so far for human gene therapy have used either MLV-A or GALV (gibbon ape leukemia 15 virus) envelopes which display some similarities (Battini, J.L., et al., J Virol. (1992) 66:1468-1475) and which use two related cell surface receptors for infection (Miller, D.G. et al., J Virol (1994) 68:8270-8276). Differences in tissuespecific expression of MLV-A or GALV receptors have been 20 reported (Kavanaugh et al., Proc Natl Acad Sci USA 91:7071-7075).

The invention will now be particularly described by way of example with reference to the accompanying drawings in which:

Figure 1.illustrates the structure and expression of CeB. The <u>env</u> gene (Xbal-Clal) of plasmid pCRIP was removed and was replaced by coinsertion of the two fragments Xbal-Sfil (restriction sites underlined) from pOXEnv and a Sfil-Clal PCR product containing the <u>bsr</u> selectable marker. This results in positioning the <u>bsr</u> start codon (shadowed) 74 bp downstream to the <u>pol</u> stop codon (bold).

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Open triangle are start codons (\underline{gaq} and \underline{bsr}), black triangles are stop codons (\underline{pol} and \underline{bsr}). The shadowed triangle is the start codon of \underline{env} , in the same reading frame with that of \underline{bsr} . SD and SA are the splice donnor and splice acceptor sites.

Figure 2 illustrates the structure and expression of FbdelPASAF.

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Immediately after the stop codon of env (bold) was inserted a non retroviral Kasl-Ncol (restriction sites underlined) linker which positions the phleo start codon (shadowed) 76 bp downstream.

Open triangle are start codons (<u>env</u> and <u>phleo</u>), black triangles are stop codons (<u>env</u> and <u>phleo</u>). SD and SA are the splice donnor and splice acceptor sites.

Figure 3 illustrates plasmids for expression of Ampho, Eco, RD114, Xeno, 10A1, GALV, VSV-G and FeLVB envelopes.

All genes are expressed in the same backbone as detailed in fig. 2. The BglII sites for ecotropic (MoMLV strain), 10A1, xenotropic (NZB.1.V6 strain) and amphotropic (4070A strain), the Ndel site of RD114 (SC3C strain, the BamHl site for both FeLVB and GALV were used as 5' ends, and linked to Mscl site immediately after the splice donor site in the leader of FB29 LTR.

Figure 4 shows the sequence of the RD114 env gene (SEQ ID No 1).

Figure 5 shows the genetic structure of gag-pol constructs.

Initiation (♥) and termination (▼) codons are shown. The thick dotted line below each construct shows MLV-derived sequences. Nucleotide positions of MLV-derived sequences are shown according to: Shinnick et al. (1981) (from nt 1 to nt 6000 with deletion of the packaging signal (DY) from Ball

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(nt 215) to PstI (nt 568), and with some further MoMLV sequences in both CeB and CeB DS- from nt 7676 to nt 7938. gag-pol and bsr genes were expressed from the same transcription unit using the either a retroviral promoter (Mo LTR) or a non retroviral promoter (hCMV) and non retroviral polyadenylation sequence (polyA). Splice donor (SD) and acceptor (SA) sites are indicated. The thin line denotes retroviral non coding sequences. The thick line shows the rabbit beta-1 globin intron B. The position of some restriction sites is indicated.

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The nucleic acid sequences of portions of constructs (as shown in Figure 5 (boxed areas)) are displayed for CeB (SEQ ID No 2, Figure 6), hCMV+intron (SEQ ID No 3, Figure 7) and hCMV+intronkaSD (SEQ ID No 4, Figure 8).

The nucleic acid sequences of portions of constructs (as shown in Figure 3 (boxed areas)) are displayed for

FbdelPASAF (SEQ ID No 5, Figure 9), FbdelPMOSAF (SEQ ID No 6, Figure 10), FbdelPGASAF (SEQ ID No 7, Figure 11),

FbdelPRDSAF (SEQ ID No8, Figure 12) and CMV10A1 (SEQ ID No 9, Figure 13) are shown.

The components of the viral particles are produced by two independent expression plasmids (gag-pol or env) which also contain selectable markers (bsr or phleo) expressed from the same transcriptional units as gag-pol or env (figs. 1& 2). The selectable markers are located downstream to gag-pol or env genes and there is an optimal distance between the stop codon of the upstream reading frames and the start codon of the selectable genes that should allow re-initiation of translation (Kozak, Mol Cell Biol. (1987) 7,:3438-3445).

Because there is no "Kozak" sequence (Kozak, Cell, (1986) 44: 283-292) required for a normal initiation of translation for

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the marker gene, they can only be expressed by re-initiation of translation after the upstream viral gene has been successfully expressed. Consequently and also because re-initiation of translation is a poorly efficient process, after transfection of these plasmids, cells resistant to the drugs corresponding to those selectable genes express high levels of the viral proteins.

To avoid viral transmission of these "helper" genomes the constructs used suitably have the classical deletions of both the packaging sequence located in the leader region and of the 3'LTR, the latter being replaced by SV40 polyadenylation sequences (Figs 1 & 2).

- Plasmid CeB is the MoMLV gag-pol-expression unit. 15 derives from pCRIP, a plasmid used to generate the constructs introduced in the CRIP and CRE packaging cell lines (Danos and Mulligan, 1988). As shown in fig. 1 for generation of plasmid CeB the env gene of pCRIP has been deleted mostly and the <u>bsr</u> selectable marker, -encoding a 20 protein conferring resistance to blasticidin (Izumi et al., Experimental Cell Research (1991) 197, 229-233) - has been inserted downstream to pol gene. There are exactly 74 bp with no ATG triplets between the stop codon of pol and the start codon of <u>bsr</u>, this allows its expression by re-25 initiation of translation on the gag-pol mRNA, after translation of the gag-pol reading frame.
- FbdelPASAF is a plasmid expressing the amphotropic env gene
 and the <u>phleo</u> selectable marker conferring resistance to
 phleomycin (Gatignol et al., FEBS Letters (1988) 230:171175). By using a PCR-mediated mutagenesis strategy which
 modifies the end of <u>env</u> gene (see fig. 2), a 76 bp linker
 was inserted between the stop codon of <u>env</u> and the start
 codon of <u>phleo</u>. This allows expression of <u>phleo</u> from the

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env mRNA by re-initiation of translation. In addition compared to known env-expressing constructs, this strategy of construction has reduced the length of sequences overlapping with the ends of conventional retroviral vectors. The env genes of Mo-MLV, FeLVB, NZB.1V6, 10A1, GALV and RD114 are expressed by plasmids FBdelPMoSAF, FBdelPBSAF, FBdelXSAF, FBdelpGSAF, FBdelp10A1SALF and FBdelPRDSAF, respectively, by using the same backbone as FBdelPASAF (fig. 3). Retroviral vectors produced with the RD114 envelope will be useful for in vivo gene delivery as comparatively to MLV ecotropic or amphotropic envelopes, virions pseudotyped with RD114 envelopes are not inactivated by human complement when they are produced by Mink Mv-1-Lu cells or by some human cells (Table 1).

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The HT1080 cell line, isolated from a human fibrosarcoma (ATCC CCL121). The TE671 cell line isolated from a human rhabdomyosarcoma (ATCC CRL 8805) (purchased from ATCC, and tested for absence of usual cell culture contaminants by ECACC), has been used for the definitive construction of packaging cell lines. HT1080 line was chosen among a panel of primate and human lines because MLV-A and RD114 efficiently rescued retroviral vectors from these cells and also because RD114 pseudotypes produced by this cell line were stable when incubated in human serum. In a standard assay (Takeuchi et al., J Virol (1994), 68, 8001-8007), these latter viruses were found more than 500 fold more stable than similar pseudotypes produced in 3T3 cells.

Another advantage for the use of non murine cells to derive packaging lines is the absence of MLV-related endogenous retroviral-like sequences (like VL30 in 3T3 cells) that can cross-package with MLV-derived retroviral vectors (Torrent et al., 1994) and generate potentially harmful recombinant retroviruses.

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The helper constructs were introduced into other cell lines (HT1080 (table 2) Mink Mv-1-Lu (table 2), 3T3 (not shown), TE671 (table 2)) for the purpose of comparisons of the efficiency of the constructs.

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As illustrated hereinafter (Table 2), the reverse transcriptase (RT) activity (provided by expression of the pol gene) in cells transfected with CeB is significantly higher than that of the same cells transfected by the parental plasmid pCRIP or that of cells chronically infected by MLV. This enhancement of viral gene expression is correlated with the titers of lacZ retroviral vectors when an envelope is provided in CeB-lacZ cells after comparison with titers of lacZ pseudotypes of either replication-competent viruses or other helper-free packaging systems.

For the generation of final packaging cell lines, the best clonal env transfectants have been selected. Packaging systems obtained in this way will be able to produce helperfree retroviral vectors at titers greater than 10⁸ infectious particles per ml, which would be 10-100 fold higher to helper-free preparations of others.

Because of the way the selectable markers are expressed (see above), growing the packaging cells in phleomycin and blasticidin selective pressure increase and stabilize the expression of the retroviral components and particularly the envelopes, as it is possible that env glycoproteins have toxic effects for the producer cells in the long term which may lead to a decrease of expression.

Such an enhancement of viral production observed with the packaging systems described herein might increase the emergence of unwanted retroviruses having recombined between the genomes of both the retroviral vector and either of the

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two packaging-deficient constructs. However, the constructs have been designed in such a way that it reduces the probability of emergence of recombinant viruses compared to the parental constructs. To check their safety, attempts have been made to detect the presence of replication-competent retroviruses by a mobilisation assay of a lacZ provirus. No RC viruses have been found in all retroviral vector preparations tested so far.

10 The following Examples illustrate the invention.

Example 1

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Preparation of Cell lines and viruses.

- The following cell lines were used:

 A204 (ATCC HTB 82), HeLa (ATCC CCL2), HT1080 (ATCC CCL121),

 MRC5 (ATCC CCL171), T24 (ATCC HTB 4), VERO (ATCC CCL81) and

 D17 (ATCC CCL183) were purchased from ATCC.
- 20 HOS, TE671 and Mv-1-Lu cells and their clones harboring MFGnlslacZ retroviral vector as described by Takeuchi et al., J Virol (1994), 68, 8001-8007.
- The above cell lines were grown in DMEM (Gibco-BRL, U.K.) supplemented with 10% fetal calf serum.

EB8 (Battini et al., J. Virol (1992) 66: 1468-1475); psiCRE, psiCRELLZ and psiCRIP (Danos et al., Proc. Natl. Acad. Sci USA (1988) 85: 6460-6464);

- Cells GP+EAM12 (Markowitz et al., Virology (1988), 167, 400-406); and
 NIH-3T3 murine fibroblasts.
- These cell lines were grown in DMEM (GIBCO-BRL, U.K.) supplemented with 10% new-born calf serum.

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Mv-1-Lu, TE671 and HT1080 cells were transfected using calcium-phosphate precipitation method (Sambrook et., "Molecular Cloning" 1989, Cold Spring Harbour Laboratory Press: New York) as described elsewhere (Battini et al., supra). CeB-transfected Mv-1-Lu, TE671 and HT1080 cells were selected with 3, 6-8 and 4 μ g/ml of blasticidin S (ICN, UK), respectively, and blasticidin-resistant colonies were isolated 2-3 weeks later. Cells transfected with the various env-expression plasmids were selected with phleomycin (CAYLA, France): 50 μ g/ml (for FBASALF-transfected cells) or 10 μ g/ml (for FBASAF-, FbdelPASAF-, FbdelPMOSAF, FBdelPIOAISAF or FBdelPRDSAF-transfected cells). Phleomycin-

FBdelPIOAISAF or FBdelPRDSAF-transfected cells). Phleomycinresistant colonies were isolated 2-3 weeks later.

Production of lacZ pseudotypes using replication competent

viruses, amphotropic murine leukemia virus (MLV-A) 1504 strain and cat endogenous virus RD114, was carried out as described previously (Takeuchi et al., J Virol (1994), 68, 8001-8007).

Example 2

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Preparation of Plasmids.

The env gene of pCRIP (Danos et al., supra) was excised by
HpaI/ClaI digestion. A 500 bp PCR-generated DNA fragment was
obtained using pSV2-bsr (Izumi et al., Experimental Cell
Research (1991), 197, 299-233) as template and a pair of
oligonucleotides:

(5'>CGGAATTCGGATCCGAGCTCGGCCCAGCCGGCCACCATGAAAACATTTAACATTTC

TC) (SEQ ID NO 2) at 5' end and
(5'>GATCCATCGATAAGCTTGGTGGTAAAACTTTT) (SEQ ID No 3) at 3'
end, with SfiI and ClaI sites, respectively. This fragment
was inserted in HpaI/ClaI sites of pCRIP by co-ligation with
a 85 bp HpaI/SfiI DNA fragment isolated from pOXEnv (Russell
et al., Nucleic Acids Research (1993), 21, 1081-1085) which

provides the end of the Moloney murine leukemia virus (MoMLV) pol gene. The resulting plasmid named CeB (Fig. 1) could express the MoMLV gag-pol gene as well as the bsr selectable marker conferring resistance to blasticidin S, both driven by the MoMLV 5'LTR promoter.

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A series of env-expression plasmids was generated using the 4070A MLV (amphotropic) env gene (Ott et al., J Virol (1990), 64, 757-766) and the FB29 Friend MLV promoter (Perryman et al., Nucleic Acid Res (1991), 19, 6950). In 10 FBASALF (Fig. 1) a BglII/ClaI fragment containing the env gene was cloned in BamHI/ClaI sites of plasmid FB3LPh which also contained the C57 Friend MLV LTR driving the expression of the phleo selection marker. A 136 bp env fragment was generated by PCR using plasmid FB3 (Heard et al., J Virol 15 (1991), 65, 4026-4032) as template and a pair of oligonucleotides: (5'>GCTCTTCGGACCCTGCATTC) (SEQ ID NO 4) at 5' end (before ClaI site) and (5'>TAGCATGGCCCCTATGGCTCGTACTCTATAGGC)(SEQ ID NO 5)at 3' end, providing a KasI restriction site immediately after the 20 env stop codon. This PCR fragment was digested using ClaI and KasI. A DNA fragment containing the FB29 LTR and the MLV-A env gene was obtained by NdeI/ClaI digestion of FBASALF. The fragments were co-ligated in Ndel/Kasl digested pUT626 (kindly provided by Daniel Drocourt, CAYLA labs, 25 France). In the resulting plasmid, named FBASAF (Fig. 1), the phleo selectable marker was expressed from the same mRNA as the env gene. A BglII restriction site was created after the MscI site at position 214 in the FB29 leader by using a commercial linker (Biolabs, France). A Ndel/BglII fragment 30 containing the FB29 LTR was co-inserted with the BglII/ClaI env fragment in NdeI/ClaI-digested FBASAF plasmid DNA, resulting in plasmid FBdelPASAF (Fig. 1). Compared to FBASAF, FBdelPASAF has a 100bp larger deletion in the leader 35 region.

Example 3

Cloning and Sequencing of the RD114 env gene The RD114 env gene was first sub-cloned in plasmid Bluescript KS+ (Stratagene) as a 3 Kb HindIII insert 5 isolated from SC3C, an RD114 infectious DNA clone (Reeves et al., J. Virol (1984), 52, 164-171). A 2.7 kb Scal-Hind III fragment of this subclone containing the RD114 env gene was sequenced (Figure 4 (SEQ ID NO 1) - EMBL accession number; X87829). The 5' non-coding sequence upstream of an NdeI site 10 was deleted by an EcoRI/NdeI digestion followed by fillingin with Klenow enzyme and self-ligation. From this plasmid, two DNA fragments were obtained: a BamHI/NcoI 2.5 Kb fragment and a 63 bp PCR-generated DNA fragment using (5'>CGCCTCATGGCCTTCATTAA) (SEQ OD NO 6) at 5' end (before 15 NotI site) and (5'>TAGCATGGCGCCTCAATCCTGAGCTTCTTCC) (SEQ ID NO 7) at 3' end, providing a KasI restriction site just after RD114 env gene stop codon. The PCR fragment was digested with NcoI and KasI. Both fragments were coinserted between BglII and KasI sites of FBdelPASAF and the 20 resulting plasmid was named FBdelPRDSAF (Fig. 1). Plasmid pCRIPAMgag- (Danos, O. et al., Proc Natl Acad Sci USA (1988) 85:6460-6464) was used for transfection.

Example 4

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Infection assays.

Target cells were seeded in 24-multiwell plates $(4 \times 10^4 \text{ cells})$ per well) and were incubated overnight. Infections were then carried out at 37°C by plating 1 ml dilutions of viral supernatants in the presence of 4 μ g/ml polybrene (Sigma) on target cells. 3h later virus-containing medium was replaced by fresh medium and infected cells were incubated for two days before X-gal staining, performed as previously described (Tailor et al., J Virol (1993), 67, 6737-6741,

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Takeuchi et al., J Virol (1994), 68, 8001-8007). Viral titers were determined by counting lacZ-positive colonies as previously described (Cosset et al., J. Virol. (1990) 64: 1070-1078). Stability of lacZ pseudotypes in fresh human serum was examined by titrating surviving virus after incubation in 1:1 mixture of virus harvest in serum-free medium and fresh human serum for 1 h at 37°C as described before (Takeuchi et al. supra).

10 Example 5

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Reverse transcriptase (RT) assay.

RT assays were performed either as described previously (Takeuchi et al. supra) or using an RT assay kit (Boehringer Mannheim, U.K.) following the manufacturer's instruction but using MnCl₂ (2 mM) instead of MgCl₂.

Example 6

20 Screening producer cell lines.

Viral particles generated with RD114 envelopes have been found to be more stable in human serum than virions with MLV-A envelopes and that the producer cell line also controls sensitivity (Takeuchi et al. supra). A panel of cell lines was screened for their ability to produce high titer viruses and for the sensitivity of these virions to human serum. To do this, cells were infected at high multiplicity with lacZ pseudotypes of either MLV-A or RD114 and cells producing helper-positive lacZ pseudotypes were established. Human HT1080 and TE671 and mink Mv-1-Lu cells were found to release high titer lacZ(RD114) and lacZ(MLV-A) viruses. LacZ(MLV-A) pseudotypes produced by HT1080 cells were more resistant to human serum than those produced by other cells. The titer of these viruses was only four-fold less following a 1 hr incubation with human serum than a

control incubation (Table 1). LacZ(RD114) pseudotypes produced by human cells or mink Mv-1-Lu cells were in general stable in human serum (Table 1). These results suggested that HT1080, TE671 and Mv-1-Lu cells provided the best combination of high lacZ titers and resistance to human serum and they were therefore used for the generation of retroviral packaging cells.

Table 1. Titer and stability of lacZ pseudotypes.

Producer	LacZ(1	LacZ(MLV-A)		LacZ(RD114) .		
cell	Titerª	Stabilityb		Stabilityb		
A204	650	<3	1,200	105		
HeLa	9 .	nd	2,000	115		
HOS	4,500	6	23,000	86		
HT1080	2,000,000	26	400,000	129		
MRC-5	450	10	1,000	nd		
T24	350	nd	1,200	nd		
TE671	15,000	2	90,000	38		
VERO	. 260	nd	90	nd		
D17	900	<1	200,000	1		
Mv-1-Lu	80,000	1	200,000	120		

a: titration on TE671 cells as lacZ i.u./ml

Example 7

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Construction of an improved gag-pol expression vector.

A MoMLV gag-pol expression plasmid, CeB (Fig. 1), was

b: % of infectivity of human serum-treated viruses compared to fetal calf serum-treated viruses

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2).

derived from pCRIP (Danos et al., Proc. Natl. Acad Aci USA (1988) 85: 6460-6464). Approximately 2 Kb of env sequence were removed from pCRIP and the bsr selectable marker, conferring resistance to blasticidin S (Izumi et al., Experimental Cell Research (1991) 197:229-233), was inserted 74 nts downstream of the gag-pol gene. This 74 nts interval had no ATG triplets and was thought to provide an optimal distance between the stop codon of the pol reading frame and the start codon of the bsr gene to allow re-initiation of translation (Kozak Mol Cell Biol., 1987, 7: 3438-3445). There was no "Kozak" consensus sequence (Kozak Cell, (1986) 44: 283-292) at the 5' end of the marker gene. Therefore, bsr could only be expressed by re-initiation of translation after the upstream gag-pol gene had been expressed. Consequently, after transfection of CeB in Mv-1-Lu/MFGnlsLacZ (ML), TE671/MFGnlsLacZ (TEL) or HT1080 cells, blasticidin S-resistant bulk populations and most cell clones expressed high levels of gag-pol proteins assessed by the reverse-transcriptase (RT) activity found in cell supernatants (Table 2). Considerably higher RT activities were found in bulk populations of CeB-transfected ML cells compared to bulk population of ML cells stably transfected with the parental pCRIP construct. Similarly the RT activities of two packaging cell lines generated using pCRIPenv- construct, psiCRE cells (Danos et al., supra) and EB8 cells (Battini supra.) were less than that of CeB transfected clones (Table 2). Finally, RT activity in CeB transfected cell supernatants was higher than that of cells chronically infected by replication-competent MLV-A (Table

Table 2. Secreted reverse transcriptase expression

Cella			_	
Cell	RT	activity ^b	LacZ	$\mathtt{Titer}^{\mathtt{c}}$

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	ML/MLV-A	1		8×104
	MLSvB	0.1		<1
	MLCRIP (bulk)	0.15	•	nd
	MLCeB (bulk)	1.7		nd
5	MLCeB1	4.2		1x10 ⁶
	MLCeB4	1.6		1x10 ⁶
	TEL/MLV-A	3.6		2x10 ⁶
	TELCeB6	5.2		4x10 ⁷
	HT1080/MLV-A	1.1		1x10 ⁶
10	HTCeB6	1.9		1x10 ⁶
	HTCeB18	2.7		2x10 ⁶
	HTCeB22 (FLY)	6.9		5x10 ⁶
	HTCeB48	5.5		3x10 ⁶
	EB8	0.22		1x10 ⁴
15	psiCRE-LLZ	1.2		1x10 ^{5d}

a: ML, Mv-1-Lu cells harboring a MFGnlslacZ provirus; TEL, TE671 cells harboring a MFGnlslacZ provirus; /MLV-A, cells chronically infected with MLV-A 1504 strain; MLSvB, ML cells transfected with a plasmid pSV2bsr alone; MLCRIP, ML cells co-transfected with pCRIP and pSV2bsr.

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b: Average of arbitrary units relative to ML/MLV-A RT activity of at least two independent experiments was shown. The standard errors did not exceed 20 % of the values.

c: titration on TE671 cells as lacZ i.u./ml. After polyclonal transfection of a plasmid which expresses MLV-A env in MLCeB clones, TELCeB clones, HTCeB clones and EB8 cells; nd, not done.

d: titration on NIH3T3 cells

To rescue infectious lacZ viruses, MLCeB and TELCeB clones
were transfected with FBASALF DNA, a plasmid designed to
express the MLV-A env gene (Fig. 1). Bulk populations of
stable FBASALF transfectants were isolated and supernatants
were titrated using TE671 cells as targets. Titers of lacZ
viruses were higher than either MLV-A infected ML or TEL

cells, or FBASALF-transfected EB8 cells (Table 2). These
data suggested that CeB was an extremely efficient MLV gagpol expression vector in mink Mv-1-Lu and TE671 cells. CeB

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was therefore used to derive packaging cells by transfection of HT1080 cells. 41/49 blasticidin S-resistant colonies had detectable levels of RT; 9 had RT activity higher than that of control MLV-A-infected HT1080 cells (data not shown). Expression of gag precursor was confirmed in cell lysates and supernatants of these 9 HTCeB clones by immunoblotting using antibodies against p30-CA (data not shown). The 4 clones with the highest expression of gag proteins (clones 6,18,22 and 48) were infected at high-multiplicity with helper free, lacZ pseudotypes bearing MLV-A envelopes (MFGnlslacZ(A)) produced by TELCeB6/FBASALF (Table 3) and then transfected with FBASALF. Supernatants of bulk, phleomycin-resistant transfectants were assessed for RT activity and lacZ titer (Table 2). Clone HTCeB22, named FLY, was found to be the best gag-pol producer clone and was used to introduce env expression vectors for the generation of packaging cell lines.

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Table 3. Titer following env construct transfection

5	Producer cell	Env source	Titerª
	psiCRIP lacZ 5	pCRIPAMgag-	6x10 ^{4b}
	GP+EAM12 lacZ 25	envAM	3x10 ^{5b}
10	TELCeB6	FBASALF° FBASAF° FbdelPASAF°	5x10 ⁷ 2x10 ⁷ 2x10 ⁷
15	TELCeB6	FBdelPASAF 1 FbdelPASAF 4 FbdelPASAF 6 FbdelPASAF 7 FbdelPASAF 8	3x10 ⁷ 2x10 ⁷ 1x10 ⁷ 5x10 ⁷
20		FbdelPASAF 8 FbdelPRDSAF 2 FbdelPRDSAF 4 FbdelPRDSAF 7 FbdelPRDSAF 8	1x10 ⁷ 1x10 ⁶ 3x10 ⁵ 1x10 ⁷ 2x10 ⁶
25	FLY ^d	FBdelPASAF 1 FbdelPASAF 4 FbdelPASAF 5 FbdelPASAF 7	1x10 ¹ 1.5x10 ⁶ 1x10 ⁶
30		FbdelPASAF 13 FbdelPASAF 14 FbdelPASAF 15 FbdelPASAF 16 FbdelPASAF 17	7x10 ⁶ 4x10 ⁶ 1x10 ⁶ 5x10 ⁶ 6x10 ⁶
35	FLYA4 lacZ 3	FBdelPRDSAF 4	2x10 ^{7b} 2.5x10 ⁶
40		FbdelPRDSAF 2 FbdelPRDSAF 6 FbdelPRDSAF 10 FbdelPRDSAF 11 FbdelPRDSAF 13 FbdelPRDSAF 17 FbdelPRDSAF 18	1x10 ⁷ 5x10 ⁶ 2x10 ⁶ 3x10 ⁶ 1x10 ⁶ 5x10 ⁶ 3x10 ⁷
45		FbdelPRDSAF 19	6x10 ⁶

Average titers of at least three independent experiments were shown. The standard errors did not exceed 30 % of the titer values.

a: titrated on TE671 cells as lacZ i.u./ml

b: results of best MFGnlslacZ producer clones.

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- c: bulk populations of env-transfectants in TELCeB6 cells.
- d: titration after bulk infection with helper-free MFGnlslacZ.

5 Example 8

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Construction of env expression vectors.

A series of MLV-A env expression plasmids were then generated (Fig. 1). In FBASALF, the env gene was inserted between two Friend-MLV LTRs, its expression driven by the 10 FB29 MLV LTR (Perryman et al., supra). Most of the packaging signal located in the leader region was deleted. This plasmid also expressed the phleo selectable marker (Gatignol et al., supra) driven by the 3' LTR. FBASAF and FBdelPASAF were then designed following the same strategy used for CeB. 15 These two vectors differed only by the extent of deletion of the packaging signal, FBdelPASAF having virtually no leader sequence. Compared to pCRIPAMgag- and pCRIPgag-2 env plasmids expressed in psiCRIP or psiCRE packaging cells (Danos et al., supra) about 5 Kb of gag-pol sequences was 20 removed. In addition the 258 bp retroviral sequence containing the end of env gene and the begining of U3 found in pCRIPAMgag- and pCRIPgag-2 was also removed. For both FBASAF and FBdelPASAF plasmids, the phleo selectable marker was inserted downstream of the env gene by positioning a 76 25 nts linker with no ATG codons between the two open-reading frames. Phleo could therefore only be expressed by reinitiation of translation by the same ribosomal unit that had expressed the upstream env open reading frame. FBdelPASAF was also used to generate FBdelPRDSAF, an RD114 30 envelope expression plasmid (Fig. 1).

After transfection of the env plasmids into TELCeB6 cells (Table 2), bulk populations of phleomycin-resistant colonies were isolated and their production of lacZ virus measured

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(Table 3). FBASALF gave a titer of $5 \times 10^7 \, \text{lacZ-i.u./ml}$, whilst titers with either FBASAF or FBdelPASAF were $2 \times 10^7 \, \text{lacZ-i.u./ml}$ (Table 3). Titers of $5 \times 10^7 \, \text{or} \, 10^7 \, \text{lacZ-i.u./ml}$ could be obtained with some FBdelPASAF cell clones or FBdelPRDSAF clones, respectively.

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As FBdelPASAF has minimal virus-derived sequences and was shown to be the safest construct (see below and Table 4), it and FBdelPRDSAF were used to generate packaging lines from FLY cells (clone HTCeB22, Table 2). Envelope expression 10 of these clones was assayed by interference to challenge with MFGnlslacZ(A) or MFGnlslacZ(RD) pseudotypes produced by TELCeB6/FBdelPASAF-7 or TELCeB6/FBdelPRDSAF-7, respectively (Table 3). The cell lines showing most interference were cross-infected at high multiplicity with these pseudotypes 15 to provide MFGnlslacZ proviruses, and supernatants were then titrated on TE671 cells (Table 3). FLY-FBdelPASAF-13 (FLYA13 packaging line) and FLY-FBdelPRDSAF-18 (FLYRD18 packaging line) gave the highest productions of lacZ viruses, around 20 107 lacZ-i.u./ml. The best MFGnlslacZ producer clones derived from either psiCRIP cells (Danos et al., supra) or GP+EAM12 cells (Markowitz et al., supra) gave approximately 50 fold lower titers (Table 3). The lacZ titers of the FLY-derived lines shown in Table 3 are lower than the best TELCeB6derived lines after transfection of either FBdelPASAF or 25 FBdelPRDSAF (Table 3). However it should be noted that the lacZ provirus expressed in TELCeB6 cells was obtained after clonal selection but was introduced polyclonally in FLYderived env-transfected cell clones. When FLY-FBdelPASAF-4 cells (FLYA4 packaging line), infected with helper-free 30 MFGnlslacZ(RD), were cloned by limiting dilution the best clones (eg. FLYA4lacZ3) were found to produce 20 times more infectious viruses than the bulk population, reaching the range of titers obtained with the best TELCeB6-FBdelPASAF 35 clones (Table 3).

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Example 9

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Assays for transfer of gag-pol or env functions.

To assay for replication-competent viruses, supernatants were used to infect TEL cells (a clone of TE671 cells harboring an MFGnlslacZ provirus). Infected cells were passaged for 6 days or longer and their supernatants were used for infection of fresh TE671 cells. No transmission of lacZ viruses could be detected (Table 4), demonstrating that the supernatants of pCRIPAMgag--, FBASALF-, FBASAF-, or FBdelPASAF-transfected TELCeB6 cells were helper-free. Similar absence of replication competent recombinant retroviruses was demonstrated using supernatant from a clone of psiCRIP-MFGnlslacZ cells or from two clones of FLYA-MFGnlslacZ cells (Table 4).

There have been reports that helper-free retroviral vector stocks may nevertheless contain recombinant retroviruses (replication incompetent) carrying either gag-pol or env genes (Bestwick et al., Proc Natl Acad Sci USA (1988), 85, 5404-5408, Cosset et al., Virology (1993), 193, 385-395, Girod et al., Virology (1995), in press). To assay for such recombinant retroviruses, mobilisation of an MFGnlslacZ provirus from two indicator cell lines which could crosscomplement potential recombinant viruses carrying either gag-pol or env functional genes was attempted. The TELCeB6 line (Table 2) expressing gag-pol proteins was used as indicator cell line to test for the presence of env recombinant (ER) viruses. The TELMOSAF indicator line expressing MoMLV env glycoproteins (obtained by transfection of FBMOSAF, a plasmid expressing the MoMLV env gene using FBASAF backbone, in TEL cells) was used to detect the presence of gag-pol recombinant retroviruses (GPR viruses). After passaging 4-8 days, the supernatants of the infected indicator cells were used to infect either human TE671 cells

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or murine NIH3T3 cells.

TELCeB6 cells transfected with various env-expressing constructs, pCRIPAMgag-, FBASAF and FBdelPASAF were compared. Although the supernatants of TELCeB6-FBdelPASAF 5 cells were devoid of replication-competent retroviruses, they were found sporadically to transfer gag-pol genomes (Table 4). No GPR viruses could be detected when less than 2x10⁵ virions were used to infect the indicator cells. Similarly TELCeB6 indicator cells infected with various 10 helper-free viruses were shown sporadically to release lacZ virions (Table 4). The number depended both on the envexpression vector used and on the virus input quantity. Compared to lacZ viruses generated using pCRIPAMgagplasmid, the frequency of detection of the env-recombinant 15 viruses was lower for supernatants generated by using FBASAF and FBdelPASAF constructs (Table 4). For FBdelPASAF construct when less than 5x10⁵ MFGnlslacZ(A) helper-free virions were used to infect the indicator cells, no ER retroviruses could be detected. From these experiments, it 20 could be estimated that a supernatant, produced from TELCeB6-FBdelPASAF cells, containing 1x107 infectious units of MFGnlslacZ retroviral vector contained no replicationcompetent virus, and about 100 gag-pol and 100 env recombinant retroviruses. 25

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Table 4. Transfer of packaging function

	Producer cell	Indicator cell	Input virus ^a	Detection ^b				
			(lacZ-i.u.)	++	+	-		
Replication competent virus								
	psiCRIP lacZ 5	TEL	2x10 ⁴	0/4	0/4	.4/		
	TELCeB6-pCRIPAMgag-	TEL	5x10 ⁶	0/4	0/4	4/4		
	TELCeB6-FBASAF	TEL	5x10 ⁶	0/4	0/4	4/4		
	TELCeB6-FBdelPASAF	TEL	5x10 ⁶	0/4	0/4	4/4		
	FLYA4 lacZ 3	TEL	$1x10^{7}$	0/4	0/4	4/4		
	FLYA4 lacZ 7	TEL	$1x10^{7}$	0/4	0/4	4/4		
Gag-pol recombinant								
	TELCeB6-FBdelPASAF 7	TELMOSAF	$2x10^{7}$	0/4	1/4	3/4		
•	TELCeB6-FBdelPASAF 7	TELMOSAF	2x10 ⁶	0/4	2/4	2/4		
•	TELCeB6-FBdelPASAF 7	TELMOSAF	2x10 ⁵	0/4	2/4	2/4		
•	TELCeB6-FBdelPASAF 7	TELMOSAF	2x10 ⁴	0/4	0/4	4/4		
		Env re	combinent		··			
•	TELCeB6-pCRIPAMgag-	TELCeB6	$5x10^{6}$	2/4	1/4	1/4		
•	TELCeB6-pCRIPAMgag-	TELCeB6	5x10 ⁵	1/4	1/4	2/4		
•	TELCeB6-pCRIPAMgag-	TELCeB6	5x10 ⁴ ·	0/4	2/4	2/4		
,	TELCeB6-FBASAF	TELCeB6	5x10 ⁶	0/4	2/4	2/4		
	TELCeB6-FBASAF	TELCeB6	5x10 ⁵	0/4	1/4	3/4		
•	TELCeB6-FBASAF	TELCeB6	5x10 ⁴	0/4	1/4	3/4		
	TELCeB6-FBdelPASAF	TELCeB6	5x10 ⁶	0/4	1/4	3/4		
	TELCeB6-FBdelPASAF	TELCeB6	5x10 ⁵	1/4	3/4	0/4		
•	TELCeB6-FBdelPASAF	TELCeB6	5x10 ⁴	0/4	0/4	4/4		

a: number of lacZ i.u. used to infect indicator cells

Titers were determined on TE671 cells for replication competent virus and env recombinant and NIH3T3 cells for

b: number of incidence out of four experiments. The ranges of lacZ titers rescued from infected indicator cells are shown for each virus input: >100 lacZ i.u./ml (++), 1-100 lacZ i.u./ml (+) and <1 lacZ i.u./ml (-).

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gag-pol recombinant.

Example 10

In order to confirm resistance to complement and absence of replication competent virus in our best packaging lines, 5 MFGnlslacZ(A) and (RD) harvested from FLYA13 and FLYRD18, respectively, after polyclonal transduction of MFGnlslacZ (Table 3 above) were tested for stability in fresh human serum and generation of replication competent virus. of MFGnlslacZ(RD) from FLYRD18 after 1 hr incubation with 3 10 independent samples of fresh human serum were 80 to 120 % of control incubations, while titers of MFGnlslacZ(A) from FLYA13 were 50 to 90 % of controls (data not shown). replication competent virus was detected in the same assay described above (Table 4) when 1 \times 10 7 i.u. each of 15 MFGnlslacZ(A) and (RD) were tested.

EXAMPLE 11.

- Generation of plasmids.

 CeB plasmid (Fig. 5) expressing MoMLV gag-pol gene, was further modified to remove the splice donor site located in the leader region. A 272 bp fragment was PCR-generated by using OUSD- (5'-TCTCGCTTCTGTTCGCGCGC) and OLSD-
- 25 (5'-TCGATCAAGCTTGCGGCCGCGGTGGTGGTGGTCGTGGTCC) as primers and further digested with BssHII and HindIII. A 1008 bp HindIII-XhoI fragment isolated from CeB (encompassing a part of leader sequence and beginning MoMLV gag) and the PCR fragment were co-inserted into pCeB from which the 1275 bp BssHII-XhoI fragment (encompassing R-U5-leader-gag) had been
- BssHII-XhoI fragment (encompassing R-U5-leader-gag) had been removed. The resulting plasmid, named pCeB DS- (Fig. 5), beared the deletion of splice donor (SD) site and a NotI restriction site created just downstream to the lost SD site.

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A series of gag-pol expression plasmids in which the MoMLV LTR promoter was replaced by the human cytomegalovirus immediate early promoter (hCMV promoter) was derived from both CeB DS- and hCMV-G (Yee et al., 1994 PNAS, 91: 9564-9568), a plasmid used as a source for the hCMV promoter. A NotI-filled/EcoRI 7260 bp fragment was isolated from CeB DS- and cloned into hCMV-G which had been opened with SalI (further rendered blunt-ended) and EcoRI to remove the VSV-G gene. The resulting plasmid was cutted with ClaI and EcoRI to remove a 1155 bp fragment encompassing sequence derived from 3'-LTR and SV40 polyA sequence and self-ligated after filling both protruding DNA ends. The resulting plasmid, named phCMV-intron (Fig. 5), had gag-pol and bsr ORFs inserted between the CMV promoter and rabbit beta-globin polyA post-transcriptional regulatory sequences.

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An intermediate plasmid was generated by sub-cloning a 7260 bp EcoRI fragment (isolated from CeB DS-) into hCMVG opened with EcoRI. A 1155 bp fragment (encompassing sequence derived from 3'-LTR and SV40 polyA sequence) was removed 20 from this intermediate plasmid which was then re-circularized by self ligation after filling both ends. The resulting plasmid, named phCMV+intron 2P (Fig. 5), was digested with NotI and the vector was treated with klenow enzyme. A 1440 bp fragment (encompassing hCMV promoter and 25 rabbit beta-1 globin intron B (Rohrbaugh et al., 1985 Mol. Cell Biol, 5: 147-160)) was isolated from phCMV+intron 2P by NotI/EcoRI digestion. This fragment was further treated with klenow enzyme and ligated back into the vector. The resulting plasmid, named hCMV+intron (Fig. 5), could express 30 gag-pol and bsr genes driven by the hCMV promoter and beared an intron sequence derived from rabbit beta-1 globin intron B having both SD and SA (splice acceptable) sites.

A 2450 bp fragment was removed from phCMV+intron 2P by

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NotI/XhoI digestion. The resulting vector fragment was then used to co-ligate a 1330 bp fragment (containing hCMV promoter + 5' end of rabbit beta-1 globin intron B (with SD site)) isolated from phCMVG by ApaI-filled/NotI digestion and a 1 kb fragment isolated from phCMV+intron 2P by NotI-filled/XhoI digestion. Compared to phCMV+intron 2P, the resulting plasmid, named hCMV+SD intron (Fig. 5), had the deletion of the 3' end of the rabbit beta-1 globin intron B and thus no SA site in the leader region.

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Construct phCMV+leader (Fig. 5) has been described elsewhere (Savard et al., unpublished). This plasmid, in which gag-pol and bsr genes were driven by the hCMV promoter, had the MoMLV SD site in the leader region.

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Gag-pol expression.

The different constructs, including the parental CeB plasmid, were analysed comparatively in a complementation assay after transfection in TEL-FBdelPASAF cells expressing 20 4070A-MLV (amphotropic) envelope and harboring a MFGnlslacZ provirus. The transient production of lacZ retroviruses as well as the stable production of lacZ retroviral vectors after selection with blasticidin S were determined (Table 5). All the constructs were able to rescue infectious lacZ retroviruses indicating the expression of gag-pol proteins 25 after transient transfection. Most likely due to the efficient hCMV and rabbit beta-1 globin intron B (post)-transcriptional regulatory sequences, hCMV+intron was particularly potent in transient retroviral vector production. However, 10 times less blasticidin-resistant 30 colonies were obtained with hCMV+intron comparatively to CeB, and stable lacZ virus production from hCMV+intron was about 5-10 times lower than that of CeB. Clonal examination of lacZ retrovirus production from blasticidin-resistant 35 colonies indicated that 80-90% of colonies could express

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high levels of gag-pol proteins for both hCMV+intron and CeB plasmids. In contrast, despite variation in their ability to form blasticidin-resistant colonies after transfection and despite their ability to express gag-pol proteins from transient transfectants, all other constructs had a weak capacity for rescuing lacZ retroviral vectors from stable transfectants (Table 5).

Table 5. Comparative study of gag-pol-bsr plasmids.

			3 3 F	TOT PERBUIT	
10	gag-pol-bsr	Transient	no clones	Stable	% gag-pol
	plasmid	(lacZ	bsr+	(lacZ	/bsr
		i.u./ml)		i.u./ml	
	Ceb	300/ml	50	10 ⁷	90%
	Ceb DS-	144/ml	5	105	50%
	hCMV+intron	ND	20	10 ⁶	50%
15	2P				
	hCMV-intron	812/ml	0	-	_
	hCMV+SD	150/ml	1000	10 ²	nd
	intron				·
:	hCMV+leader	328/ml	1000	10 ² -10 ³	nd
20	hCMV+intron	12000/ml	5	106-107	80왕 :

Northern blot analyses were performed on stable transfectants (blasticidin-resistant) obtained with some of the gag-pol-bsr plasmids. As expected, the results (not shown) displayed a correlation between expression of gag-pol mRNAs and gag-pol protein expression detected by rescue analysis (Table 5). CeB construct was found to produce 2-3 fold more gag-pol mRNAs compared to hCMV+intron.

Interestingly, an unexpected 2.45 kb RNA band was found for hCMV+intron construct at a ratio of 2:1 compared to the abundancy of the gag-pol mRNA band (at 5.95 kb). Further

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investigations by using other probes revealed that a cryptic splice donnor (SD) site located in the gag gene (right in the middle of the CA coding region at position 1596-1597 -numbering according to Shinnick et al., 1981 Nature (London) 293: 543-548) was activated in this latter construct. The 2.45 RNA species, lacking the 3' half of the gag gene and most of the pol gene, is unlikely to give rise to any useful translational product. It is therefore interesting to notice that hCMV+intron construct was able to give rise to slightly more transcripts (gag-pol 5.95 mRNA + 2.45 alternative RNA band) compared to gag-pol mRNA expressed from CeB construct. Therefore we decided to inactivate the cryptic SD site in the hCMV+intron construct in order to increase the ratio of gag-pol mRNAs.

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Assays for transfer of gag-pol functions.

Although the supernatants of pacakaging cell lines generated with CeB gag-pol expression contruct were devoid of replication-competent retroviruses, they were found sporadically to transfer gag-pol genomes (example 9, Table 4) (Cosset et al., 1995 J. Virol 69: 7430-7436). Because gag-pol-bsr constructs generated here by using the hCMV promoter had much less retroviral sequences homologous to the retroviral vector than the parental CeB construct (Fig. 5), they are less likely to give rise to gag-pol recombinant (GPR) viruses. Therefore, the most efficient gag-pol-bsr plasmids, hCMV+intron and CeB, were further analysed for emergence of GPR viruses. To assay for such recombinant retroviruses, we attempted to mobilise an lacZ provirus from an indicator cell lines which could cross-complement potential recombinant viruses carrying gag-pol functional genes. Results displayed in Table 6 showed that consistently with data reported previously (example 9, Table 4) (Cosset et al., 1995 Supra), lacZ retrovirus vectors generated by using CeB gag-pol construct were contaminated with GPR viruses. In

contrast lacZ retrovirus vectors generated by using hCMV+intron construct were completely devoid of such GPR viruses, suggesting that this construct was improved compared to CeB with respects with emergence of recombinant viruses.

Table 6. Comparative study of gag-pol-bsr plasmids.

plasmid	input virus (lacZ i.u.)ª	no of experiments giving titres ofb		
СеВ	5x10 ⁶	5	3	0
	5x10 ⁵	2	4	2
	5x10 ⁴	0	1	7
hCMV+intron	5x10 ⁶	0	0	8
	5x10 ⁵	0	0	8
	5x10 ⁴	0	0	8

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4x10E4 cells of TEL/MOSAF in 24 wells were challenged with lacZ(A) of i.u. indicated in the table (a), and incubated at 37°C for 3 days. Cells were trypsinized and transferred into small flasks. Cell sup was harvested on day 5 after lacZ(A) challenge and plated on either TE571 (not shown) and 3T3 cells (b). No lacZ was mobilized into TE671 at all. LacZ(A) from CMV-int 10 again did not rescue lacZ from TEL/MOSAF.

Example 12

Generic primers to detect D-type (Medstrand and Blomberg 25 30

J. Virol. (1993) 67:6778-6787) , C-type (Shih et al., J Virol. (1989) 63:64-75), human endogenous virus RTVL-H (Wilkinson et al., J.Virol. (1993) 67:2981-2989), by RT-PCR were employed (Patience et al., supra). Primers to detect mouse endogenous VL30 element (Adams et al Mol.Cel.Biol. (1988) 8:2989-2998), and MFGnlslacZ RNA were designed and synthesized (TABLE X). Overnight supernatants (in 4ml of culture medium) from 106 cells of GP+EAM12lacZ25, FLYA4lacZ3

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and TELCeB6FBASALF cells (Table 3) were harvested and centrifuged in sucrose gradient as described previously (Patience et al., J.Virol., 70:2654-2657). Fractions containing retrovirus particles were collected, and RNA extracted. One twentieth of the RNA preparation or dilution's thereof were applied to RT-PCR as described previously (Table X). A 1/200 of RNA harvested from GP+EAM12lacZ25 cells was positive for VL30 RNA. MFGnlslacZ RNA was found from 1/20 of RNA from GP+EAM12lacZ and TELCeB6FBASALF cells and 1/200 of RNA from FLYA4lacZ3 cells. The primer combinations for RTVL-H, C- and D-type RNA did not give detectable PCR product.

Table 7. RT-PCR detection of endogenous retrovirus RNA associated with virus particles.

			rt-pcr of v	irion associ	ated RNA fromª
20	RNA	primer (5'-3') forward(F)/reverse(R)	GP+EAM12 lacZ25	FLYA4 lacZ3	TELCeB6F BASALF
25	MFGnls lacZ	F) CTCTGGCTCACAGTACGACGTAR) CCATCAATCCGGTAGGTTTTCC		· ++	+
30	C-type	F) CARRGKTTCAARAACWSYCCCAR) AGYARVGTAGCNGGGTTHAGG	AC -	· -	-
	D-type	F) TCCCCTTGGAATACTCCTGTTTR) CATTCCTTGTGGTAAAACTTTC		-	-
35	RTVL-H	F) CCTCACCCTGATCACRYTTG R) GAATTATGTCTGACAGAAGGG	ТИ	-	· <u>-</u>
	VL30	F) GTTGACATCTGCAGAGAAAGAC R) TCTGAGGTCTGTACACACAATG		NT	NT

a:-,not detected; + detected in 1/20 RNA preparation; ++ detected in 1/200 RNA preparation; NT, not tested because the cells do not possess the corresponding genes.

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EXAMPLE 13.

Generation of gag-pol pre-packaging cells by using TE671 cells.

10 CeB, a plasmid designed to over-express MoMLV gag and pol proteins was introduced in TE671 human rhabdomyosarcoma cells (ATCC CRL8805). After selection with blasticidin, 50 bsr-positive colonies were isolated and the RT (reverse transcriptase) activity was analysed in their supernatants.

12 TE671-CeB (TECeB) clones with high RT activity were selected for further analysis. The best TECeB clone, clone #15, had a RT activity roughly equivalent to that TELCeB6 cells (Cosset et al., J. Virol. 69:7430-7436 (1995); see

- also Example 7, Table 6 in this patent application) but displayed 2-3 fold more gag-precursors into cells as demonstrated in immunoblots by using anti-CA antibodies. The biological activity of gag-pol proteins expressed in the six best TECeB clones was further confirmed by their ability to produce infectious retroviruses in a complementation assay.
- A lacZ provirus was introduced into each of the TECeB clones by polyclonal cross-infection by using lacZ(RD114) helper-free retrovirus vectors. FBMOSALF, a MoMLV env expression plasmid (Cosset et al., J. Virol. 69:6314-6322), was then transfected in each of the TECeB-lacZ lines and in the TELCeB6 cell line for comparison. After solvetion with
- TELCeB6 cell line for comparison. After selection with phleomycin, the titer of lacZ retrovirus vectors was determined in the supernantant of pools of phleomycin-resistant colonies for each TECEB-lacZ-FBMOSALF lines. A

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good correlation was found between gag-pol expression into the TE-CeB clones (as determined by RT-assays and anti-gag immunoblots) and their ability to release infectious lacZ particles. TE-CeB15 cells could release approximately the same number of lacZ particles when compared to TELCeB6 cells although TELCeB6 cells had the advantage of being selected for lacZ expression (Cosset et al., J. Virol. 69:7430-7436 (1995)). TE-CeB15 cells were therefore used to derive retroviral packaging cell lines.

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Construction of env-expression plasmids.

A series of plasmid (Fig. 3) was designed to allow expression of different retroviral envelope genes (isolated from MoMLV, GALV -Gibbon Ape Leukemia Virus-, and MLV-10A1). FBdelPMOSAF (Fig. 3, nucleotide sequence in Fig. 10) and 15 FBdelP10A1SAF, expressing ecotropic MoMLV or MLV-10A1 envelopes, were generated by replacing the BglII/ClaI fragment from FBdelPASAF (Cosset et al., J. Virol. 69:7430-7436 (1995); see also Example 7, Fig. 2 and nucleotide sequence in Fig. 9) encompassing most of the env gene and 20 splice acceptor site with that of MoMLV (position 5407 to 7679, Shinnik et al., 1981) or with that of MLV-10A1 (Ott et al., J. Virol. 64:757-766 (1990)). Nucleotides 7514-7516 of GALV (Delassus et al., Virology 173:205-213 (1989)) were mutated by PCR-mediated mutagenesis 25 to create a ClaI site (AAG to CGA), thereby introducing a conservative modification (a lysine (amino-acid 665 of GALV env precursor) to an arginine). The BamHI/ClaI fragment (nts 4994 (Delassus et al. Virology 173:205-213 (1989)) to 7517)

was then sub-cloned into FBdelPASAF in which the BglII/ClaI encompassing most of the env gene and splice acceptor site had been removed. The resulting plasmid, expressing GALV

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envelope glycoproteins, was named FBdelPGASAF (Fig. 3, nucleotide sequence in Fig. 11).

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CMV10A1 was generated by inserting a Klenow enzyme-filled EagI/SalI fragment from FBdelP10A1SAF (encompassing 10A1 MLV env gene and phleo selectable marker) into hCMV-G digested with BamHI and filled with Klenow enzyme. The resulting plasmid, CMV10A1 (Fig. 3 and nucleotide sequence in Fig. 13) could express 10A1 envelopes under control of the hCMV promoter and the phleo selectable marker by translation reinitiation.

Generation of a multi-tropic set of TE671-based retroviral packaging lines.

FBdelPRDSAF (Fig. 3, nucleotide sequence in Fig. 12),

FBdelPASAF, FBdelPGASAF, FBdelPMOSAF and FBdelP10A1SAF were independently introduced into cells of the TE-CeB15 prepackaging line, expressing MoMLV gag-pol proteins.

Transfected cells were phleomycin-selected and 15-20 phleoresistant colonies were isolated for each env-expression plasmid transfected.

Individual colonies were the second of the TE-CeB15 prepackaging line, expressing MoMLV gag-pol proteins.

Individual colonies were then analysed for expression of envelope glycoproteins by immunoblots on cell lysates by using antibodies against RD114 SU glycoproteins or against Rausher leukemia virus SU (to screen MoMLV, MLV-4070A and MLV-10A1 env-producer clones) or against GALV. The best env-producer colonies as determined in this assay were further analysed by a complementation assay after introducing a lacZ retroviral vector. LacZ pseudotypes released from the different packaging cell lines were titrated by using NIH 3T3 cells or TE671 cells as target. Titers higher than 1x107 lacZ i.u./ml were obtained for the best clones. Depending on the envelope specificities expressed in these cells, the new

TE671-based retroviral packaging cell lines were named TE-FLYE, TE-FLYA, TE-FLYRD, TE-FLY10A1, and TE-FLYGA and could express the MoMLV, MLV-4070A, RD114, MLV-10A1, and GALV env genes, respectively.

Assays for detecting replication-competent retroviruses (RCRs) were performed in the supernatants of these cells and were negative (less than 1/ml).

TE671 cells are very potent for transient expression resulting in more than 95% of cells expressing transgene 10 three days after plasmid transfection (Hatziioannou and Cosset, unpublished data, (1996)). The ability of retroviral packaging cell lines to transiently produce retroviral vectors is of crucial importance for gene therapy where vectors carrying toxic gene have to be prepared. Transient 15 expression of retroviral vectors was comparatively determined from cells of the TE-FLYA line and from the BING line (Pear et al., Proc Natl Acad Sci U S A 90, 8392-6 (1993)), a retroviral packaging cell line designed to transiently express retroviral vectors. Results (Table 8) 20 showed that TE-FLYA cells were more efficient for transient expression of a lacZ retroviral vector hence resulting in higher titers.

Table 8. Comparative study of transient production of lacZ vectors.

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packaging cell line	cell numberª	% transfected cells ^b	transient titer°
BING	281	5.3	2x10 ²
TE-FLYA	117	35	1.3x10 ³

Cells were transfected by MFGnlslacZ retroviral vectors with calcium phosphate precipitation method and titers of of lacZ vectors (c) released in cell

supernatant were determined as lacZ i.u./ml at day 3 following transfection. The relative number of cells (a) (average per microscope field) and the % of transfected cells (b) determined after X-gal staining are shown.

Retroviral vectors prepared from TE671-based packaging cell lines were analysed for their sensitivity to human—complement mediated inactivation. Experiments were conducted as previously described (Cosset et al., J. Virol. 69:7430—7436 (1995); see also Example 10 in this patent application) by using three human sera of individual donnors (Table 9). As expected MLV-A prepared from mouse 3T3 cells were highly sensitive to inactivation after 1 hr incubation with sera. In contrast, titers of lacZ vectors produced from TE-FLYRD cells were 17 to 55% of control incubations, while titers of lacZ vectors from TE-FLYA cells were 1 to 30% of controls.

Table 9. Human serum sensitivity of viruses produced from TE671-based packaging cell lines.

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Virus from:	hu56ª	hu57°	BTSª
3T3/A	<0.2, <0.2	<0.2, <0.2	<0.2, <0.2
TE-FLYE .	15, 7.8	16, 11	48, 60
TE-FLYA	1, 0.6	2.2, 7.1	28, 19
TE-FLYRD	17, 22	30, 44	54, 63

Three human fresh serum samples were tested in duplicate; hu56 (A+), hu57(AB+), BTS(AB+). (a) % control (average for FCS and opti-MEM treatment) is shown.

CLAIMS:

- 1. A recombinant expression vector comprising a gene of interest and a selectable marker gene, wherein the selectable marker gene is arranged downstream of the gene of interest and a stop codon associated with the gene of interest is spaced from a start codon of said selectable marker gene at a distance which is sufficient to ensure that said selectable marker protein is expressed from the corresponding mRNA as a result of translation reinitiation.
- 2. A recombinant expression vector according to claim 1 wherein the vector is a viral vector.
- 3. A recombinant expression vector according to claim 2 wherein the vector is a retroviral vector.
- 4. A recombinant expression vector according to any one of claims 1 to 3 wherein the gene of interest is included as part of a viral packaging construct.
- 5. A recombinant expression vector according to any one of the preceding claims wherein the number of nucleotides in the space between the stop codon of the gene of interest and the start codon of the selectable marker is in the range of from 20 to 200 nucleotides.
- 6. A recombinant expression vector according to claim 5 wherein the number of nucleotides in the space between the stop codon of the gene of interest and the start codon of the selectable marker is in the range of from 60 to 80 nucleotides.
- 7. A process for producing a cell line in which a gene of interest is expressed, which process comprises: transforming host cells with an expression vector

- according to any one of the claims 1 to 6; and selectable those cells where expression of the selection marker gene may be detected.
- 8. A process according to claim 7 wherein the host cell is a eukaryotic cell.
- 9. A host cell transformed with a recombinant expression vector according to any one of the claims 1 to 6.
- A retroviral packaging cell line comprising a host 10. cell transformed with a first and a second recombinant expression vector, said first recombinant expression vector having a packaging-deficient construct comprising a viral gag-pol gene and a first selectable marker gene downstream thereof, and said second recombinant expression vector having a packagingdeficient construct comprising a viral env gene and a second selectable marker gene downstream thereof; wherein the start codon of the first and second selectable markers are spaced from the stop codons of the viral gag-pol gene and the viral env gene respectively by a distance which ensures that said selectable marker protein is expressed from the corresponding mRNA as a result of translation reinitiation.
- 11. A retroviral packaging cell line according to claim 10 wherein the first selectable marker is a bsr selectable marker and the second selectable marker is a phleo selectable marker.
- 12. A retroviral packaging cell line according to any one of claims 10 or 11 wherein the packaging-deficient construct comprising the viral gag-pol gene and first selectable marker is the CeB (SEQ ID No 2) expression construct.

- 13. A retroviral packaging cell line according to any one of claims 10 or 11 wherein the packaging-deficient construct comprising the viral env gene and second selectable marker is the FBdelPASAF (SEQ ID No 5), the FBdelPMOSAF (SEQ ID No 6), the FbdelPGASAF (SEQ ID No 7), the FbdelPRDSAF (SEQ ID No 8), the FbdelPXSAF (Fig. 3), the FbdelP10A1SAF (Fig. 3), or the FBdelPVSVGSAF (Fig. 3) expression construct.
- 14. A retroviral packaging cell line according to any one of claims 10 or 11 wherein the recombinant expression vector is a packaging-deficient retroviral helper construct.
- 15. A retroviral packaging cell line according to claim 14 wherein the overlapping sequences between the genomes of the retroviral vector and the packaging-deficient construct is reduced by minimizing the extent of non-coding retroviral sequences in the packaging-deficient genome.
- 16. A retroviral packaging cell line according to any one of claims 10 to 15 wherein the viral gag-pol gene and the selectable marker are expressed under the control of a non-retroviral promoter.
- 17. A retroviral packaging cell line according to claim 16 wherein the promoter is fused to rabbit beta-1 globin intron.
- 18. A retroviral packaging cell line according to claim 16 or claim 17 wherein the promoter is a hCMV promoter.
- 19. A retroviral packaging cell line according to any one of claims 16 to claim 18 wherein the viral gag-pol gene and the selectable marker is a hCMV+intron (SEQ

- ID No3) or a hCMV+intronkaSD (SEQ ID No 4) expression construct.
- 20. A retroviral packaging cell line according to anyone of claims 10 to 15 wherein the viral env gene and the selectable marker are under the control of a non-retroviral promoter.
- 21. A retroviral packaging cell line according to claim 20 wherein the promoter is fused to rabbit beta-1 globin intron.
- 22. A retroviral packaging cell line according to claim 20 or claim 21 wherein the promoter is a hCMV promoter.
- 23. A retroviral packaging cell line according any one of claims 20 to 22 wherein the viral env gene and the selectable marker is a CMV10A1 (SEQ ID No 9) expression construct.
- 24. A retroviral packaging cell line according to any one of claims 10 to 23 wherein the cell line is the HT1080 line, the TE671 line, the 3T3 line, the 293 line or the MV-1-1U line.
- 25. A retroviral packaging cell line according to anyone of claims 10 to 24 wherein the retroviral packaging cells comprises human HT1080 cells and express RD114 envelopes.
- 26. A retroviral packaging cell line according to anyone of claims 10 to 24 wherein the retroviral packaging cells comprises human TE671 cells and express RD114 envelopes.

27. A process for producing a retroviral packaging cell line in which a gene of interest in expressed, which process comprises:

transforming host cells with a first and a second recombinant expression vector, said first recombinant expression having a packaging-deficient vector construct comprising a viral gag-pol gene and a first selectable marker gene downstream thereof, and said recombinant expression vector having packaging-deficient construct comprising a viral env gene and a second selectable marker gene downstream thereof; wherein the start codon of the first and second selectable markers are spaced from the stop codons of the viral gag-pol gene and the viral env gene respectively by a distance which ensures that said selectable marker protein is expressed from the corresponding mRNA as a result of translation reinitiation; and

selecting transformed cells which express said first and/or second marker genes.

- 28. A packaging deficient construct for use in a process according to claim 27, which expresses a viral gag-pol gene and a selectable marker wherein a start codon of the selectable marker is spaced from a stop codon of the viral gag-pol gene by a distance which ensures that said selectable marker protein is expressed from the corresponding mRNA as a result of translation reinitiation.
- 29. A packaging deficient construct for use in a process according to claim 27, which expresses a viral env gene and a selectable marker gene; wherein a start codon of the selectable marker is spaced from a stop codon of the viral env gene by a distance which ensures that said selectable marker protein is

expressed from the corresponding mRNA as a result of translation reinitiation.

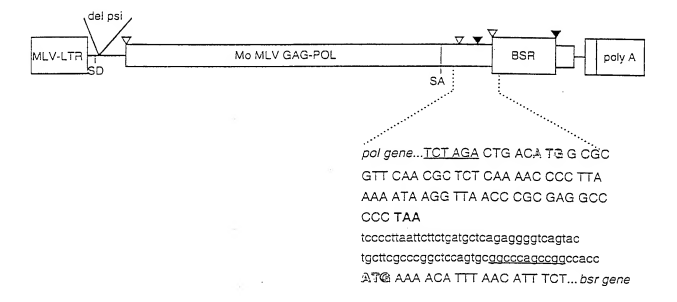


Figure 1. Schematic structure of CeB expression vector

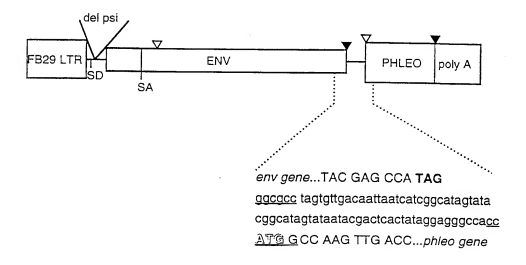


Figure 2. Schematic structure of FBdelPASF expression vector

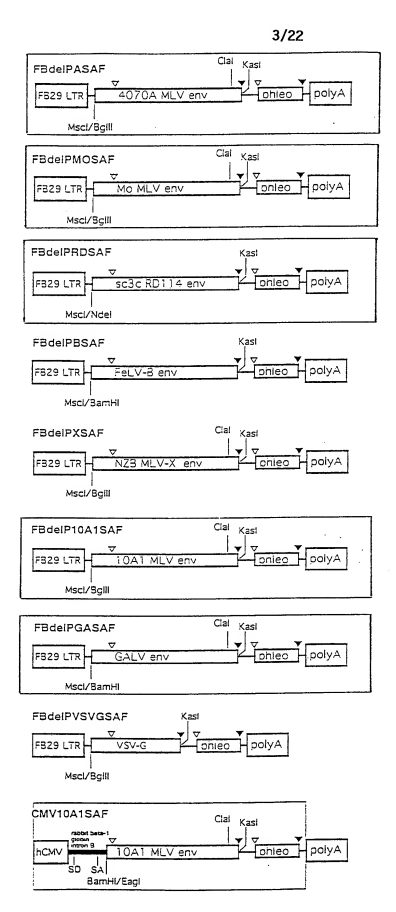


Figure 3. Schematic structure of env expression vectors

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NGAGCTCAGGACAGGTAGAAAGAATGAATAGAACAATAAAAGAGACCCTTACTAAATTGA 60 CCTTAGAGACTGGCTTAAAAGATTGGAGACGCCTCCTATCTCTGGCTTTGTTAAGAGCCA 120 GAAATACGCCCAACCGTTTTCGGCTCACCCCATATGAAATCCTTTATGGGGGACCCCCC 180 CTTTGTCAACCTTGCTCAATTCCTTCTCCCCCTCCGATCCTAAGACTGATTTACAAGCCC 240 GACTAAAAGGGCTGCAAGGCCTGCAGGCCCAAATCTGGACACCCCTGGCCGAATTGTACC 300 GGCCAGGACATCCACAAACTAGCCACCCATTTCAGGTGGGAGACTCCGTGTACGTCCGGC 360 GGCACCGCTCTCAAGGATTGGAGCCTCGTTGGAAGGGACCTTACATCGTCCTGCTGACCA 420 CGCCCACCGCCATAAAGGTTGACGGGATCGCCGCCTGGATTCACGCATCGCACGCCAAGG 480 CAGCCCCAAAAACCCCTGGACCAGAAACTCCCAAAACCTGGAAGCTCCGCCGTTCGGAGA 540 ACCCTCTTAAGATAAGACTCTCCCGTGTCTGACTGACTCACCTTGTCCCTGTACTAA 600 CCCAAAATGAAACTCCCAACAGGAATGGTCATTTTATGTAGCCTAATAATAGTTCGGGCA 660 GGGTTTGACGACCCCCGCAAGGCTATCGCATTAGTACAAAAACAACATGGTAAACCATGC 720 CCAGGCAAGACGGCCTACTTAATGACCAACCAAAAATGGAAATGCAGAGTCACTCCAAAA 840 ATCTCACCTAGCGGGGGAGAACTCCAGAACTGCCCCTGTAACACTTTCCAGGACTCGATG 900 CACAGTTCTTGTTATACTGAATACCGGCAATGCAGGCGAATTAATAAGACATACTACACG 960 GCCACCTTGCTTAAAATACGGTCTGGGAGCCTCAACGAGGTACAGATATTACAAAACCCC 1020 AATCAGCTCCTACAGTCCCCTTGTAGGGGCTCTATAAATCAGCCCGTTTGCTGGAGTGCC 1080 ACAGCCCCCATCCATATCTCCGATGGTGGAGGACCCCTCGATACTAAGAGAGTGTGGACA 1140 GTCCAAAAAAGGCTAGAACAAATTCATAAGGCTATGACTCCTGAACTTCAATACCACCCC 1200 TTAGCCCTGCCCAAAGTCAGAGATGACCTTAGCCTTGATGCACGGACTTTTGATATCCTG 1260 AATACCACTTTTAGGTTACTCCAGATGTCCAATTTTAGCCTTGCCCAAGATTGTTGGCTC 1320 TGTTTAAAACTAGGTACCCCTACCCCTCTTGCGATACCCACTCCCTCTTTAACCTACTCC 1380 CTAGCAGACTCCCTAGCGAATGCCTCCTGTCAGATTATACCTCCCCTCTTGGTTCAACCG 1440 ATGCAGTTCTCCAACTCGTCCTGTTTATCTTCCCCTTTCATTAACGATACGGAACAAATA 1500 GACTTAGGTGCAGTCACCTTTACTAACTGCACCTCTGTAGCCAATGTCAGTAGTCCTTTA 1560 TGTGCCCTAAACGGGTCAGTCTTCCTCTGTGGAAATAACATGGCATACACCTATTTACCC 1620 CAAAACTGGACCAGACTTTGCGTCCAAGCCTCCTCCTCCCGACATTGACATCAACCCG 1680 GGGGATGAGCCAGTCCCCATTCCTGCCATTGATCATTATACATAGACCTAAACGAGCT 1740 GTACAGTTCATCCCTTTACTAGCTGGACTGGGAATCACCGCAGCATTCACCACCGGAGCT ACAGGCCTAGGTGTCTCCGTCACCCAGTATACAAAATTATCCCATCAGTTAATATCTGAT 1860 GTCCAAGTCTTATCCGGTACCATACAAGATTTACAAGACCAGGTAGACTCGTTAGCTGAA 1920 GTAGTTCTCCAAAATAGGAGGGGACTGGACCTACTAACGGCAGAACAAGGAGGAATTTGT 1980 TTAGCCTTACAAGAAAATGCTGTTTTTATGCTAACAAGTCAGGAATTGTGAGAAACAAA 2040 TGGACCGGGCTGCAGGGCTTTCTTCCGTACCTCCTACCTCCTGGGACCCCTACTCACC 2160 CTCCTACTCATACTAACCATTGGGCCATGCGTTTTCAGTCGCCTCATGGCCTTCATTAAT 2220 GATAGACTTAATGTTGTACATGCCATGGTGCTGGCCCAGCAATACCAAGCACTCAAAGCT 2280 GAGGAAGAAGCTCAGGATTGAGCTTCCGGGACAAAAGCAGGGGGGAATGAGAAGTCAGAA 2340 CCCCCCACCTTTGCTACATAAATAACCGCTTTCATTTCGCTTCTGTAAAACGCTTATGCG 2400 CCCCACCTAGCCGGAAAGTCCCCAGCCGCTACGCAACCCGGGCCCCGAGTTGCATCAGC 2460

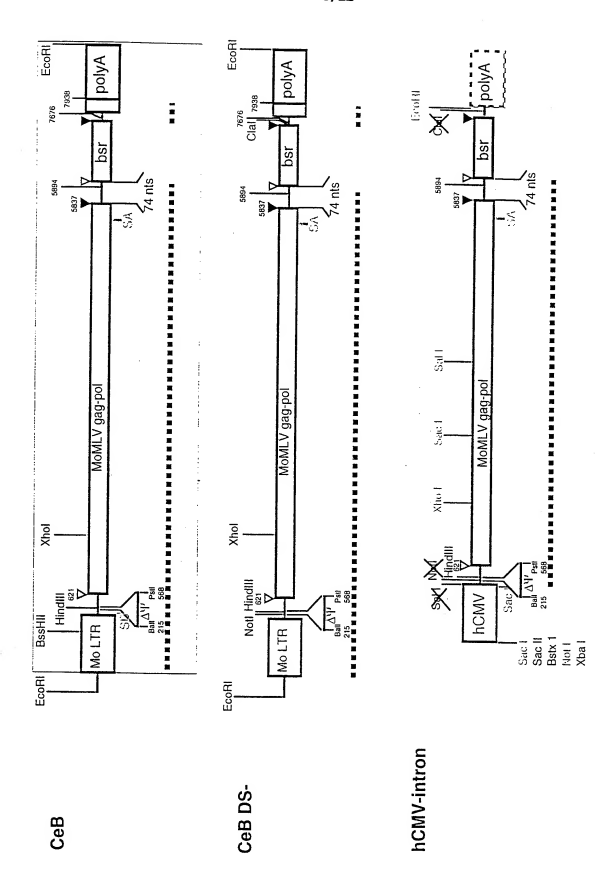


Figure 5. Genetic structure of gag-pol constructs (page 1/3)

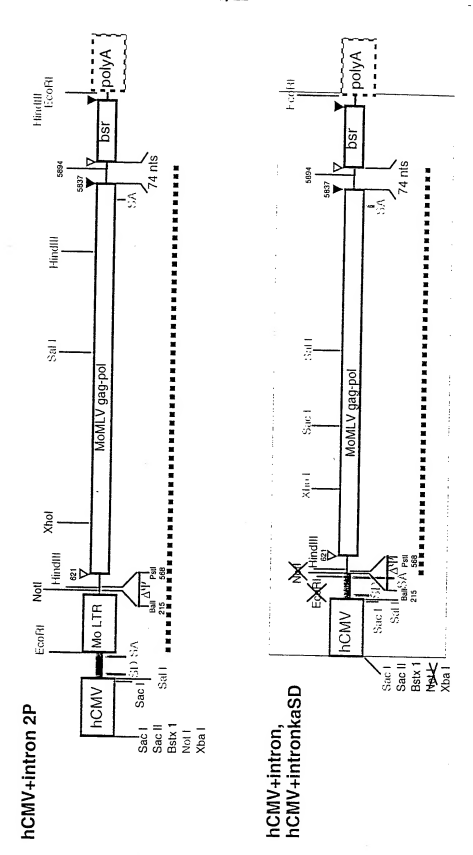


Figure 5. Genetic structure of gag-pol constructs (page 2/3)



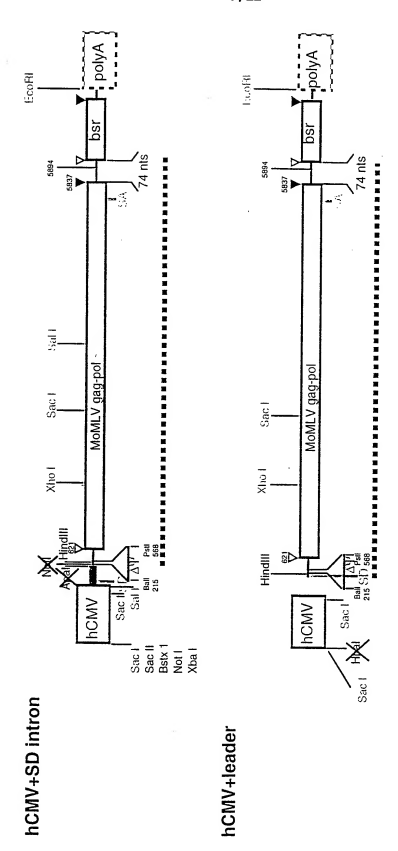


Figure 5. Genetic structure of gag-pol constructs (page 3/3)

Figure 6. CeB Sequence 8/22

AATGAAAGAC	CCCACCTGTA	GGTTTGGCAA	GCTAGCTTAA	GTAACGCCAT	TTTGCAAGGC	- 60
	ACATAACTGA					120
AGCTGAATAT	GGGCCAAACA	GGATATCTGT	GGTAAGCAGT	TCCTGCCCCG	GCTCAGGGCC	180
	GGAACAGCTG					240
	GGGCCAAGAA					300
	CAGATGTTTC					360
TGAACTAACC	AATCAGTTCG	CTTCTCGCTT	CTGTTCGCGC	GCTTCTGCTC	CCCGAGCTCA	420
ATAAAAGAGC	CCACAACCCC	TCACTCGGGG	CGCCAGTCCT	CCGATTGACT	GAGTCGCCCG	480
	TATCCAATAA					540
	TCTCCTCTGA					600
CTCGTCCGGG	ATCGGGAGAC	CCCTGCCCAG	GGACCACCGA	CCCACCACCG	GGAGGTAAGC	660
TGGAAGCTTC	TGCAGCATCG	TTCTGTGTTG	TCTCTGTCTG	ACTGTGTTTC	TGTATTTGTC	720
	GGCCAGACTG					780
TGTCGAGCGG	ATCGCTCACA	ACCAGTCGGT	AGATGTCAAG	AAGAGACGTT	GGGTTACCTT	840
CTGCTCTGCA	GAATGGCCAA	CCTTTAACGT	CGGATGGCCG	CGAGACGGCA	CCTTTAACCG	900
AGACCTCATC	ACCCAGGTTA	AGATCAAGGT	CTTTTCACCT	GGCCCGCATG	GACACCCAGA	960
CCAGGTCCCC	TACATCGTGA	CCTGGGAAGC	CTTGGCTTTT	GACCCCCCTC	CCTGGGTCAA	1020
GCCCTTTGTA	CACCCTAAGC	CTCCGCCTCC	TCTTCCTCCA	TCCGCCCCGT	CTCTCCCCCT	1080
TGAACCTCCT	CGTTCGACCC	CGCCTCGATC	CTCCCTTTAT	CCAGCCCTCA	CTCCTTCTCT	1140
AGGCGCCAAA	CCTAAACCTC	AAGTTCTTTC	TGACAGTGGG	GGGCCGCTCA	TCGACCTACT	1200
	CCCCCGCCTT					1260
	GCGACCCCTG					1320
	CGGGAGCCCC					1380
	AACGGACAGC					1440
	AACCCTTCTT					1500
	ACCCATCAGC					1560
	GAAAAACAAC					1620
	ACTCAACTGC					1680
	ACCACCCAGG					1740
	CAAAACGCGG					1800
	AATGAGTCTC					1860
	TATGACCCTG					1920
	GCCCCAGACA					1980
	GATTTGGTTA					2040
	GAACGTATCA					2100
	AAAGAGAAAG					2160
	GTTAGTGGAC					2220
	GACCAGTGTG					2280
	CGAGGACCTC					2340
	CAGGGTCAGG					2400
	ACCTTCCTGG					2460
	AGTGATAAGT					2520
	GATCGCAAAG					2580
	TGTCCCTATC					2640
	GAGGGATCAG					2700
GTTGACCCTA	AATATAGAAG	ATGAGCATCG	GCTACATGAG	ACCTCAAAAG	AGCCAGATGT	2760
	TCCACATGGC					2820
	GTTCGCCAAG					2880
	CAATACCCCA				CCCACATACA	2940
	AAACCAGGGA					
	GAAGACATCC					3060
					TCTGCCTGAG	3120 3180
					AGATGGGAAT	
					CCACCCTGTT	
					ACTTGATCCT	
					GCCAACAAGG	
TACTCGGGCC	CTCTTACAAA	CCCTACCCAA	CCTCCCCTAT	CGGGCCTCGG	CCAAGAAAGC	3480
CCAAATTTCC	CAGAAACACC	TCD ACTAGGAA	CCCCTATCT	CTANAGACA	GTCAGAGATG	3540
					CCCCTCGACA	
ACTAAGGGAG	TTCCTAGGGA	CCCCACCCOTO	CTGTCGCCTC	TCGATCCCTC	GGTTTGCAGA	3660
					GGGGCCCAGA	
					CCCTGGGGTT	
					ACGCCAAAGG	
					CCAAAAAGCT	
					TTGCCGTACT	
	GCAGGCAAGC					4020
					GGATGACTCA	
						2000

Figure 6. CeB Sequence 9/22 2

CTATCAGGCC	TTGCTTTTGG	ACACGGACCG	GGTCCAGTTC	GGACCGGTGG	TAGCCCTGAA	4140
CCCGGCTACG	CTGCTCCCAC	TGCCTGAGGA	AGGGCTGCAA	CACAACTGCC	TTGATATCCT	4200
GGCCGAAGCC	CACGGAACCC	GACCCGACCT	AACGGACCAG	CCGCTCCCAG	ACGCCGACCA	4260
CACCTGGTAC	ACGGATGGAA	GCAGTCTCTT	ACAAGAGGGA	CAGCGTAAGG	CGGGAGCTGC	4320
GGTGACCACC	GAGACCGAGG	TAATCTGGGC	TAAAGCCCTG	CCAGCCGGGA	CATCCGCTCA	4380
GCGGGCTGAA	CTGATAGCAC	TCACCCAGGC	CCTAAAGATG	GCAGAAGGTA	AGAAGCTAAA	
TGTTTATACT	GATAGCCGTT	ATGCTTTTGC	TACTGCCCAT	ATCCATGGAG		4440
AAGGCGTGGG	TTGCTCACAT	CAGAAGGCAA	AGAGATCAAA	AATAAAGACG	ACAMCMMCCCC	4500
CCTACTAAAA	GCCCTCTTTC	TGCCCAAAAG	ACTTAGCATA	ATTCCATTCTC	CACCACAGO	4560
AAAGGGACAC	AGCGCCGAGG	CTAGAGGCAA	CCGCATGCCT	GACCAACCCC	CAGGACATCA	4620
AGCCATCACA	GAGACTCCAG	ACACCTCTAC	CCGCATGGCT	CAAAAAMMCAM	CCCGAAAGGC	4680
CTCAGAACAT	TTTCATTACA	CACTCACTCA	TATA A ACCAC	CONTRACT	CACCCTACAC	4740
TTATCATAAA	ACAAAGAAGT	AMMCCCMCMA	CCAACCAAAA	CIAACCAAGT	TGGGGGCCAT	4800
T	TTATTAGACT	MITGGGICIA	CCHAGGAAAA	CCTGTGATGC	CTGACCAGTT	4860
CCCTCTCCTA	CACACARGACT	TTCTTCATCA	GCTGACTCAC	CTCAGCTTCT	CAAAAATGAA	4920
AAAMAMCACM	GAGAGAAGCC	ACAGTCCCTA	CTACATGCTG	AACCGGGATC	GAACACTCAA	4980
AGACCCAACT	GAGACCTGCA	AAGCTTGTGC	ACAAGTCAAC	GCCAGCAAGT	CTGCCGTTAA	5040
CAMAAACCC	AGGGTCCGCG	GGCATCGGCC	CGGCACTCAT	TGGGAGATCG	ATTTCACCGA	5100
GATAAAGCCC	GGATTGTATG	GCTATAAATA	TCTTCTAGTT	TTTATAGATA	CCTTTTCTGG	5160
CIGGATAGAA	GCCTTCCCAA	CCAAGAAAGA	AACCGCCAAG	GTCGTAACCA	AGAAGCTACT	5220
AGAGGAGATC	TTCCCCAGGT	TCGGCATGCC	TCAGGTATTG	GGAACTGACA	ATGGGCCTGC	5280
CTTCGTCTCC	AAGGTGAGTC	AGACAGTGGC	CGATCTGTTG	GGGATTGATT	GGAAATTACA	5340
TTGTGCATAC	AGACCCCAAA	GCTCAGGCCA	GGTAGAAAGA	ATGAATAGAA	CCATCAAGGA	5400
GACTTTAACT	AAATTAACGC	TTGCAACTGG	CTCTAGAGAC	TGGGTGCTCC	TACTCCCCTT	5460
AGCCCTGTAC	CGAGCCCGCA	ACACGCCGGG	CCCCCATGGC	CTCACCCCAT	ATGAGATCTT	5520
ATATGGGGCA	CCCCCGCCCC	TTGTAAACTT	CCCTGACCCT	GACATGACAA	GAGTTACTAA	5580
CAGCCCCTCT	CTCCAAGCTC	ACTTACAGGC	TCTCTACTTA	GTCCAGCACG	AAGTCTGGAG	5640
ACCTCTGGCG	GCAGCCTACC	AAGAACAACT	GGACCGACCG	GTGGTACCTC	ACCCTTACCG	5700
AGTCGGCGAC	ACAGTGTGGG	TCCGCCGACA	CCAGACTAAG	AACCTAGAAC	CTCGCTGGA A	5760
AGGACCTTAC	ACAGTCCTGC	TGACCACCCC	CACCGCCCTC	AAAGTAGACG	GCATCGCAGC	5820
TTGGATACAC	GCCGCCCACG	TGAAGGCTGC	CGACCCCGGG	GGTGGACCAT	CCTCTAGACT	5880
GACATGGCGC	GTTCAACGCT	CTCAAAACCC	CTTAAAAATA	AGGTTAACCC	GCGAGGCCCC	5940
CTAATCCCCT	TAATTCTTCT	GATGCTCAGA	GGGGTCAGTA	CTGCTTCGCC	CGGCTCCAGT	6000
GCGGCCCAGC	CGGCCACCAT	GAAAACATTT	AACATTTCTC	AACAAGATCT	AGAATTAGTA	6060
GAAGTAGCGA	CAGAGAAGAT	TACAATGCTT	TATGAGGATA	ATAAACATCA	TGTGGGAGCG	6120
GCAATTCGTA	CGAAAACAGG	AGAAATCATT	TCGGCAGTAC	ATATTGAAGC	GTATATAGGA	6180
CGAGTAACTG	TTTGTGCAGA	AGCCATTGCG	ATTGGTAGTG	CAGTTTCGAA	TGGACAAAAG	6240
GATTTTGACA	CGATTGTAGC	TGTTAGACAC	CCTTATTCTG	ACGAAGTAGA	TAGAAGTATT	6300
CGAGTGGTAA	GTCCTTGTGG	TATGTGTAGG	GAGTTGATTT	CAGACTATGC	ACCAGATTCT	6360
TTTGTGTTAA	TAGAAATGAA	TGGCAAGTTA	GTCAAAACTA	CGATTGAAGA	ACTCATTCCA	
CTCAAATATA	CCCGAAATTA	AAAGTTTTAC	CACCAAGCTT	ATCGATTAGT	CCAATICCA	6420
AAAGACAGGA	TATCAGTGGT	CCAGGCTCTA	CTTTTTTCACTC	ALCANTATION	CCACITIGIT	6480
CCTATAGAGT	ACGAGCCATA	GATAAAATAA		TTAGTCTCCA		6540
GGAATGAAAG	ACCCCACCTG	TAGGTTTGGC	AACCTACCTT	AAGTAACGCC	AUDURCCAAC	6600
GCATGGAAAA	ATACATAACT	GAGAATAGAG		CAAGGTCAGG		6660
ACAGTCGAGA	ACTTGTTTAT	TGCAGCTTAT	AATCCTTACA	AAMAAACCAA	MACAGAIGGA MACAGAIGGA	6720
AATTTCACAA	ATAAAGCATT	TOCAGCITAT	CATTCTACT	CTCCTTTCCTC	CAAACTCACA	6780
AATGTATCTT	ATCATGTCTG	GATCCCCACC	AACCTCCTCTT	CECECCECA	AAACTCATC	6840
CTCCTCTACT	TGAGAGGACA	TTCCC A ATTCAT	AGGCTCCTCT	MCCA CCCMCM	AAACCCTAAC	6900
GTTAATTAGG	TCACTTAACA	AAAACCAAAM	TCCCTACCCCA	TCCACCCTCT	ACCCONT	6960
AAGGGTAATT	TTAAAATATC	TCCCAACTCC	CUMCCACACA	TITITCACAG	ACCGCTTTCT	7020
AAACAGCCCA	CAAATGTCAA	CAGCACAAAC	ATACAACCTCC	TCACCOMMOCO	AAGTGTTGGT	7080
AACACCCTGC	TCATCAAGAA	CCACTCTCCT	TACAAGCIG	CMA A MCMCCC	ACAAGGGCCC	7140
CACATTTTCC	CCACCTGTGT	ACCEPTOR A A	AMAMCMACMC	GTAATGTGCA	AAACAGGAGG	7200
AGGAACCCAG	CACTCCACTG	CAMAACCAMM	ATATCTAGTG	A A A A CA COCC	TACTTGGATC	7260
GTTCATCTCC	TGACTGTCAA	CMCMACCATT	ATCCTTATCC	AAAACAGCCT	TGTGGTCAGT	7320
GGTCCTGC	TOACIGICAA	CACCORCAN	TTTTGGGGTTT	ACAGTTTGAG	CAGGATATTT	7380
ATCAAAATT	TTTGCTAACA	MCCCMMMMCC	A COA COA TOTA	TUCUCACCAA	CAGCAAAAA	7440
TCAATCCAAC	GACCCTTGAA	CACHERACCO	AGCACCATTT	TCATGAGTTT	TTTGTGTCCC	7500
CCACAMCAAA	TTTAACATAG	CAGTTACCCC	AATAACCTCA	GTTTTTAACAG	TAACAGCTTC	7560
CONCATCAMA	ATATTTCCAC	AGGTTAAGTC	CTCATTTAAA	TTAGGCAAAG	GAATTC	7616

Figure 7. hCMV+intron Sequence

AGATCTCCCG	ATCCCCTATG	GTCGACTCTC	AGTACAATCT	GCTCTGATGC	CGCATAGTTA	- 60
AGCCAGTATC	TGCTCCCTGC	ттстстсттс	GAGGTCGCTC	AGTAGTGCCC	GAGCAAAATT	
TAAGCTACAA	CAAGGCAAGG	CTTGACCGAC	A A TOTO CATCA	ACA A MCMCCCC	TAGGGTTAGG	
CCTTTTTCCCC	TCCTTCCCCA	mcma coccac	AMITGUMIGM	AGAATCIGCI	TAGGGTTAGG	180
A COULT TO COC	ACUDULIUUL A	TGTACGGGCC	AGATATACGC	GTTGACATTG	ATTATTGACT	240
AGITATTAAT	AGTAATCAAT	TACGGGGTCA	. TTAGTTCATA	. GCCCATATAT	GGAGTTCCGC	300
GTTACATAAC	TTACGGTAAA	TGGCCCGCCT	GGCTGACCGC	CCAACGACCC	CCGCCCATTG	2.50
ACGTCAATAA	. TGACGTATGT	TCCCATAGTA	. ACGCCAATAG	GGACTTTCCA	THEACHTEAA	400
TGGGTGGACT	ATTTACGGTA	. AACTGCCCAC	TTGGCAGTAC	ATCAACTCTA	$\Phi C \lambda \Phi \lambda \Phi C C C \lambda$	
AGTACGCCCC	CTATTGACGT	CAATGACGGT	AAATGGCCCG	CCTCCCATTA	TCAIAIGCCA	
ATGACCTTAT	GGGACTTTCC	TACTOCCOAC	ma Camema co	MITAJOUTJJ	TGCCCAGTAC	540
ATCCTCATCC	CCTTTTTCC	CMACAMCAAM	COCCOCCOCC	TATTAGTCAT	CGCTATTACC	600
MMMCC33CMC	GGTTTTGGCA	GTACATCAAT	GGGCGTGGAT	AGCGGTTTGA	CTCACGGGGA	660
TITCCAAGIC	TCCACCCCAT	TGACGTCAAT	GGGAGTTTGT	TTTGGCACCA	AAATCAACGG	720
GACTTTCCAA	AATGTCGTAA	CAACTCCGCC	CCATTGACGC	AAATGGGCGG	TACCCCTCTA	780
CGGTGGGAGG	TCTATATAAG	CAGAGCTCTC	TGGCTAACTA	GAGAACCCAC	TO COURT A COLO	840
GCTTATCGAA	ATGTCGACTG	AGAACTTCAG	GGTGAGTTTG	GGGACCCTTG	Δ $\overline{\mathbf{u}}$ \mathbf{u}	
CTTTTTCGCT	ATTGTAAAAT	TCATGTTATA	TGGAGGGGGC	AAAGTTTTCA	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	900
TAGAATGGGA	AGATGTCCCT	TCTATCACCA	TCCA CCCTCA	MCAMA A TOTAL	GGGTGTTGTT	960
CTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TCTTCACAAC	CAMMONMON	TOGACCCICA	TGATAATTTT	GTTTCTTTCA	1020
TTCTTCTC	TGTTGACAAC	CATTGTCTCC	TCTTATTTC	TTTTCATTTT	CTGTAACTTT	1080
1 COTTAAAC	TTTAGCTTGC	ATTTGTAACG	AATTTTTAAA	TTCACTTTTG	TTTATTTGTC	1140
AGATTGTAAG	TACTTTCTCT	AATCACTTTT	TTTTCAAGGC	AATCAGGGTA	TATTATATTG	1200
TACTTCAGCA	CAGTTTTAGA	GAACAATTGT	TATAATTAAA	TGATAAGGTA	$C_{\Delta} \Delta T_{\Delta} T_{\Delta} T_{\Delta} T_{\Delta} T_{\Delta} T_{\Delta}$	1260
GCATATAAAT	TCTGGCTGGC	GTGGAAATAT	TCTTATTGGT	AGAAACAACT	$\Delta C \Delta T C C T C C T$	
CATCATCCTG	CCTTTCTCTT	ТАТССТТАСА	ΑΤΓΑΤΑΤΑΤΑ	CTCTTTCACA	MCACCAMAAA	1320
ATACTCTGAG	TCCAAACCGG	GCCCCTCTCC	TIL OTTITITION	CAUCCOURGE	IGAGGATAAA	1380
ACAGCTCCTC	CCCAACCTCC	#CC##C##C#	TAACCATGIT	CATGCCTTCT	TCTTTTTCCT	1440
CCAACCHHCH	GGCAACGTGC	TGGTTGTTGT	GCTGTCTCAT	CATTTTGGCA	AGAATTGGCC	1500
GCAAGCTTCT	GCAGCATCGT	TCTGTGTTGT	CTCTGTCTGA	CTGTGTTTCT	GTATTTGTCT	1560
GAGAATATGG	GCCAGACTGT	TACCACTCCC	TTAAGTTTGA	CCTTAGGTCA	CTGGAAAGAT	1620
GTCGAGCGGA	TCGCTCACAA	CCAGTCGGTA	GATGTCAAGA	AGAGACGTTG	CCTTACCTTC	1680
TGCTCTGCAG	AATGGCCAAC	CTTTAACGTC	GGATGGCCGC	GAGACGCCAC	COTTACCITC	
GACCTCATCA	CCCAGGTTAA	GATCAAGGTC	THE TOTAL COLUMN	GCCCCCARCC	2C1CCCACCCA	1740
CAGGTCCCCT	ACATCGTGAC	CTCCCAACCC	TTTTCTCTG	ACCCCCCCTCC	ACACCCAGAC	1800
CCCTTTCTAC	ACCCUAACCC	TICCCCCTCCT	COMMONTAL	ACCCCCCTCC	CTGGGTCAAG	1860
CAACCTCCTC	ACCCTAAGCC	TCCGCCTCCT	CTTCCTCCAT	CCGCCCCGTC	TCTCCCCCTT	1920
CCCCCCAAAC	GTTCGACCCC	GCCTCGATCC	TCCCTTTATC	CAGCCCTCAC	TCCTTCTCTA	1980
3G2GCCAAAC	CTAAACCTCA	AGTTCTTTCT	GACAGTGGGG	GGCCGCTCAT	CGACCTACTT	2040
ACAGAAGACC	CCCCGCCTTA	TAGGGACCCA	AGACCACCCC	CTTCCGACAG	GGACGGAAAT	2100
GGTGGAGAAG	CGACCCCTGC	GGGAGAGGCA	CCGGACCCCT	CCCCAATGGC	ATCTCGCCTA	2160
CGTGGGAGAC	GGGAGCCCCC	TGTGGCCGAC	TCCACTACCT	CGCAGGCATT	CCCCCTCCCC	2220
GCAGGAGGAA	ACGGACAGCT	TCAATACTGG	CCGTTCTCCT	CTTCTCACCT	TTACAACTCC	
AAAAATAATA	ACCCTTCTTT	ΤΤΟΤΟΣΑΙΟΣΤ	CCACCTAAAC	TCACACCTC	CAMCCAACIGG	2280
GTTCTCATCA	CCCATCAGCC	CACCOCCCAC	CACTOTIANAC	1GACAGCICI	GATCGAGTCT	2340
ACCGGAGAAG	CCCATCAGCC	CACCIGGGAC	GACTGTCAGC	AGCTGTTGGG	GACTCTGCTG	2400
CCCCCCCCC	AAAAACAACG	GGTGCTCTTA	GAGGCTAGAA	AGGCGGTGCG	GGGCGATGAT	2460
TOGGCGCCCCA	CTCAACTGCC	CAATGAAGTC	GATGCCGCTT	TTCCCCTCGA	GCGCCCAGAC	2520
TGGGATTACA	CCACCCAGGC	AGGTAGGAAC	CACCTAGTCC	ACTATCGCCA	GTTGCTCCTA	2580
GCGGGTCTCC	AAAACGCGGG	CAGAAGCCCC	ACCAATTTGG	CCAAGGTAAA	AGGAATAACA	2640
CAAGGGCCCA	ATGAGTCTCC	CTCGGCCTTC	CTAGAGAGAC	TTAAGGAAGC	CTATCGCAGG	2700
TACACTCCTT	ATGACCCTGA	GGACCCAGGG	CAAGAAACTA	ATCTCTCTAT	CTCTTTCGCAGG	
TGGCAGTCTG	CCCCAGACAT	TGGGAGAAG	TTACACACCT	TACAACAMM	CICILICATI	2760
ACCCTTCGAG	ATTTGGTTAG	ACACCCACAA	TINGMOMONI	IMGMAGATTT	AAAAAACAAG	2820
GAAAGAGAGG	A A C C M A M C A C	CACACAAA	AAGATCTTTA	ATAAACGAGA	AACCCCGGAA	2880
CAMCACCACA	AACGTATCAG	GAGAGAAACA	GAGGAAAAAG	AAGAACGCCG	TAGGACAGAG	2940
CCCACECEC	AAGAGAAAGA	AAGAGATCGT	AGGAGACATA	GAGAGATGAG	CAAGCTATTG	3000
GCCACTGTCG	TTAGTGGACA	GAAACAGGAT	AGACAGGGAG	GAGAACGAAG	GAGGTCCCAA	3060
CTCGATCGCG	ACCAGTGTGC	CTACTGCAAA	GAAAAGGGGC	ACTGGGCTAA	AGATTGTCCC	3120
AAGAAACCAC	GAGGACCTCG	GGGACCAAGA	CCCCAGACCT	CCCTCCTGAC	ССТАСАТСАС	3180
TAGGGAGGTC	AGGGTCAGGA	GCCCCCCCT	GAACCCAGGA	TAACCCTCAA	AGTCCCCCC	
CAACCCGTCA	CCTTCCTGGT	AGATACTGGG	GCCCAACACT	CCCTCCTCAC	CCAAAATCCT	3240
GGACCCCTAA	GTGATAAGTC	TCCCTCCCTC	CAACCCCCTA	CMCCACCAAA	CCAAAATCCT	
TGGACCACCC	ATCCCANACE	1GCC1GGG1C	CAAGGGGCTA	CTGGAGGAAA	GCGGTATCGC	3360
DDJAJJADDI	ATCGCAAAGT	ACATCTAGCT	ACCGGTAAGG	TCACCCACTC	TTTCCTCCAT	3420
GIACCAGACT	GTCCCTATCC	TCTGTTAGGA	AGAGATTTGC	TGACTAAACT	AAAAGCCCAA	3480
ATCCACTTTG	AGGGATCAGG	AGCTCAGGTT	ATGGGACCAA	TGGGGCAGCC-	CCTGCAAGTG	3540
'I''I'GACCCTAA	ATATAGAAGA	TGAGCATCGG	CTACATGAGA	CCTCAAAAGA	GCCAGATGTT	3600
TCTCTAGGGT	CCACATGGCT	GTCTGATTTT	CCTCAGGCCT	GGGCGGAAAC	CGGGGGCATG	3660
GGACTGGCAG	TTCGCCAAGC	TCCTCTGATC	ATACCTCTGA	AAGCAACCTC	TACCCCCGTG	
TCCATAAAAC	AATACCCCAT	GTCACAAGAA	CCCAGACTCC	CCAMCAACCIC	CCACATACAG	
AGACTICTUTCC	ACCACCCAI	A COCCOUR CCC	CCCAGACTGG	GGATCAAGCC	CCACATACAG	
CCCCumbyycz	ACCAGGGAAT	ACTGGTACCC	TGCCAGTCCC	CCTGGAACAC	GCCCCTGCTA	3840
CCCGIIMAGA	AACCAGGAC	TAATGATTAT	AGGCCTGTCC	AGGATCTGAG	AGAAGTCAAC	3900
AAGCGGGTGG	AAGACATCCA	CCCCACCGTG	CCCAACCCTT	ACAACCTCTT	CACCCCCCTC	2060
CCACCGTCCC	ACCAGTGGTA	CACTGTGCTT	GATTTAAAGG	AUCCCUUTUTUT	CTCCCTCACA	4020
CTCCACCCCA	CCAGTCAGCC	TCTCTTCGCC	TTTGAGTGGA	GAGATCCAGA	GATGGGAATC	4080
						4000

Figure 7. hCMV+intron Sequence

TCAGGACAAT	TGACCTGGAC	CAGACTCCCA	CAGGGTTTCA	AAAACAGTCC	CACCCTGTTT	4140
GATGAGGCAC	TGCACAGAGA	CCTAGCAGAC	TTCCGGATCC	AGCACCCAGA	CTTCATCCTC	4200
CTACAGTACG	TGGATGACTT	ACTGCTGGCC	GCCACTTCTG	AGCTAGACTG	CCAACAACCM	4260
ACTOGGGCCC	TGTTACAAAC	CCTAGGGAAC	CTCGGGTATC	GGGCCTCGGC	CAAGAAACCC	4320
CAAATTTGCC	AGAAACAGGT	CAAGTATCTG	GGGTATCTTC	TAAAAGAGGG	TCAGAGATGG	4200
CTGACTGAGG	CCAGAAAAGA	GACTGTGATG	GGGCAGCCTA	CTCCGAAGAC	CCCTCGAGAA	4440
CTAAGGGAGT	TCCTAGGGAC	GGCAGGCTTC	TGTCGCCTCT	GGATCCCTCG	CTTTCCACAA	4500
ATGGCAGCCC	CCTTGTACCC	TCTCACCAAA	ACGGGGACTC	ጥርጥጥጥ ል ልጥጥር	GGGCCCACAC	
CAACAAAAGG	CCTATCAAGA	AATCAAGCAA	GCTCTTCTAA	CTGCCCCAGC	CCTCCCCTTC	4560
CCAGATTTGA	CTAAGCCCTT	TGAACTCTTT	GTCGACGAGA	AGCAGGGCTA	CCCCAAACCM	4620
GTCCTAACGC	AAAAACTGGG	ACCTTGGCGT	CGGCCGGTGG	CCTACCTCTC	COCCAAAGGI	4680
GACCCAGTAG	CAGCTGGGTG	GCCCCCTTGC	CTACGGATGG	TAGCAGCCAT	TCCCCCTA CTC	4740
ACAAAGGATG	CAGGCAAGCT	AACCATGGGA	CAGCCACTAG	TCATTCTCC	CCCCCAMCCA	4800
GTAGAGGCAC	TAGTCAAACA	ACCCCCGAC	CCCTCCCTTT	CCAACCCCCC	CCCCCATGCA	
TATCAGGCCT	TGCTTTTGGA	CACGGACCGG	GTCCAGTTCG	GACCCCCCC	ACCOMMO	4920
CCGGCTACGC	TGCTCCCACT	GCCTGAGGAA	GGGCTGCAAC	ACA ACTICCOM	MCAMAMORRO	4980
GCCGAAGCCC	ACGGAACCCG	ACCCGACCTA	ACGGACCAGC	CCCTCCCACA	CCCCCACCAC	5040
ACCTGGTACA	CGGATGGAAG	CAGTCTCTTA	CAAGAGGGAC	ACCCURA ACCC	CGCCGACCAC	
GTGACCACCG	AGACCGAGGT	AATCTCCCCT	A A ACCCCTCC	CACCCCCAC	AGGGAGCTGCG	5160
CGGGCTGAAC	TGATAGCACT	CACCCAGGCC	CTANACCTIC	CAGCCGGAC	ATCCGCTCAG	5220
GTTTATACTG	ATAGCCGTTA	TCCTTTTCCT	ACTICCCCATA	TCCATCCACA	GAAGCTAAAT	5280
AGGCGTGGGT	TGCTCACATC	AGAACCCAAA	CACAMCAAAA	1CCATGGAGA	AATATACAGA	5340
CTACTAAAAC	CCCTCTTTCT	CCCCAAA	GAGATCAAAA	ATAAAGACGA	GATCTTGGCC	5400
AAGGGACACA	GCGCCCACCC	MACACCCA AC	CTTAGCATAA	TCCATTGTCC	AGGACATCAA	5460
GCCATCACAG	GCGCCGAGGC	CACCECTAC	CGGATGGCTG	ACCAAGCGGC	CCGAAAGGCA	5520
TCAGAACATT	AGACTCCAGA	CACCTCTACC	CTCCTCATAG	AAAATTCATC	ACCCTACACC	5580
TATCATATA	TTCATTACAC	AGTGACTGAT	ATAAAGGACC	TAACCAAGTT	GGGGGCCATT	5640
ስርጥጥጥጥር እ አጥ	CAAAGAAGTA	TIGGGICTAC	CAAGGAAAAC	CTGTGATGCC	TGACCAGTTT	5700
CCTCTCCTAC	TATTAGACTT	TCTTCATCAG	CTGACTCACC	TCAGCTTCTC	AAAAATGAAG	5760
AATATCACTC	AGAGAAGCCA	CAGTCCCTAC	TACATGCTGA	ACCGGGATCG	AACACTCAAA	5820
CACCCAACTA	AGACCTGCAA	AGCTTGTGCA	CAAGTCAACG	CCAGCAAGTC	TGCCGTTAAA	5880
ATAAACCCCC	GGGTCCGCGG	GCATCGGCCC	GGCACTCATT	GGGAGATCGA	TTTCACCGAG	5940
TEGATACAAC	GATTGTATGG	CTATAAATAT	CTTCTAGTTT	TTATAGATAC	CTTTTCTGGC	6000
CACCACARCR	CCTTCCCAAC	CAAGAAAGAA	ACCGCCAAGG	TCGTAACCAA	GAAGCTACTA	6060
TTCCTCTCTC	TCCCCAGGTT	CGGCATGCCT	CAGGTATTGG	GAACTGACAA	TGGGCCTGCC	6120
TCTCCATACA	AGGTGAGTCA	GACAGTGGCC	GATCTGTTGG	GGATTGATTG	GAAATTACAT	6180
A COMMON A CONA	GACCCCAAAG	CTCAGGCCAG	GTAGAAAGAA	TGAATAGAAC	CATCAAGGAG	6240
CCCCCCTACC	AATTAACGCT	TGCAACTGGC	TCTAGAGACT	GGGTGCTCCT	ACTCCCCTTA	6300
TATCCCCCAC	GAGCCCGCAA	CACGCCGGGC	CCCCATGGCC	TCACCCCATA	TGAGATCTTA	6360
ACCCCCTCTC	CCCCGCCCCT	TGTAAACTTC	CCTGACCCTG	ACATGACAAG	AGTTACTAAC	6420
CCTCTCTC	TCCAAGCTCA	CTTACAGGCT	CTCTACTTAG	TCCAGCACGA	AGTCTGGAGA	6480
CCTCTGGCGG	CAGCCTACCA	AGAACAACTG	GACCGACCGG	TGGTACCTCA	CCCTTACCGA	6540
GCACCTTACA	CAGTGTGGGT	CCGCCGACAC	CAGACTAAGA	ACCTAGAACC	TCGCTGGAAA	6600
TCCATACACC	CAGTCCTGCT	GACCACCCC	ACCGCCCTCA	AAGTAGACGG	CATCGCAGCT	6660
ACATECECEC	CCGCCCACGT	GAAGGCTGCC	GACCCCGGGG	GTGGACCATC	CTCTAGACTG	6720
TA ATTCCCCTT	TTCAACGCTC	TCAAAACCCC	TTAAAAATAA	GGTTAACCCG	CGAGGCCCCC	6780
CCCCCCTT	AATTCTTCTG	ATGCTCAGAG	GGGTCAGTAC	TGCTTCGCCC	GGCTCCAGTG	6840
AACTACCCAC	GGCCACCATG	AAAACATTTA	ACATTTCTCA	ACAAGATCTA	GAATTAGTAG	6900
CAAMMCCMAC	AGAGAAGATT	ACAATGCTTT	ATGAGGATAA	TAAACATCAT	GTGGGAGCGG	6960
CAGTAC	GAAAACAGGA	GAAATCATTT	CGGCAGTACA	TATTGAAGCG	TATATAGGAC	7020
AMERICA CAC	TTGTGCAGAA	GCCATTGCGA	TTGGTAGTGC	AGTTTCGAAT	GGACAAAAGG	7080
CACTCCCTAAC	GATTGTAGCT	GTTAGACACC	CTTATTCTGA	CGAAGTAGAT	AGAAGTATTC	7140
THE CHECKER A.T.	TCCTTGTGGT	ATGTGTAGGG	AGTTGATTTC	AGACTATGCA	CCAGATTGTT	7200
TIGIGITAAT	AGAAATGAAT	GGCAAGTTAG	TCAAAACTAC	GATTGAAGAA	CTCATTCCAC	7260
ICAAATATAC	CCGAAATTAA	AAGTTTTACC	ACCAAGCTTA	TCGAATTC		7308

Figure 8. hCMV+intronkaSD Sequence 1

AGATCTCCCG	ATCCCCTATG	GTCGACTCTC	AGTACAATCT	CCTCTGATGC	CCCATACTTA	60
A C C C A C M A M C	TCCTCCCTCC	mmamamamma	ar comecome	7.077.070.00	COCHIAGIIA	-
MGCCMGIMIC	TGCTCCCTGC	TTGTGTGTTG	GAGGTCGCTG	AGTAGTGCGC	GAGCAAAATT	120
TAAGCTACAA	CAAGGCAAGG	CTTGACCGAC	AATTGCATGA	AGAATCTGCT	TAGGGTTAGG	100
CCMMMMCCCCC	TCCTTCCCTTCC	CITOMCCOMC	MITOCHION	HUMAICIGCI	LAGGGIIAGG	180
CGTTTTGCGC	TGCTTCGCGA	TGTACGGGCC	AGATATACGC	GTTGACATTG	ATTATTGACT	240
Αςττατταατ	AGTAATCAAT	TACCCCCTCA	$mm \times mm \subset X \times X$	CCCCAMAMAM	CCACMMOGGG	
	THE THIT CAME	IACGGGGICA	IINGIICAIA	GCCCATATAT	GGAGTTCCGC	300
GTTACATAAC	TTACGGTAAA	TGGCCCGCCT	GGCTGACCGC	CCAACGACCC	CCGCCCATTC	360
አሮሮሞሮጳ አመጻ አ	mc x ccm x mcm	mccca ma cma	3.000003.3.03.0	GG3 CMMMCG3		
ACGICAAIAA	TGACGTATGT	TCCCATAGTA	ACGCCAATAG	GGACTTTCCA	TTGACGTCAA	420
TGGGTGGACT	ATTTACGGTA	AACTCCCCAC	TTCCCACTAC	ልጥር እ እርጥር መ አ	mcamamacaa	
10001001101	TITTICOOTA	AACIGCCCAC	TIGGCAGIAC	AICAAGIGIA	ICATATGCCA	480
AGTACGCCCC	CTATTGACGT	CAATGACGGT	AAATGGCCCG	CCTGGCATTA	TCCCCACTAC	540
A MC A CCMM A M	CCCACMMMCC	ma comecaca c	ma ca moma co	#3.##3.##a	TOCCCHOIAC	
AIGACCITAI	GGGACTTTCC	TACTTGGCAG	TACATCTACG	TATTAGTCAT	CGCTATTACC	600
ATGGTGATGC	GGTTTTGGCA	GTACATCAAT	GCCCCTCCAT	ACCCCTTTCA	CTCACCCCCA	
	55111160671	OTTICITE CENT	COCCIOCAL	MOCGGIIIGA	ADDDDJAJIJ	660
TTTCCAAGTC	TCCACCCCAT	TGACGTCAAT	GGGAGTTTGT	TTTGGCACCA	AAATCAACGG	720
CACTTTCCAA	AATGTCGTAA	CAACTCCCCC	CCAMMCACCC	A A A MCCCCCCC	m1.00000	
GACTITCCAA	AAIGICGIAA	CAACICCGCC	CCALIGACGC	AAATGGGCGG	TAGGCGTGTA	780
CGGTGGGAGG	TCTATATAAG	CAGAGCTCTC	TGGCTAACTA	GAGAACCCAC	ጥርርጥጥ አ አ ርጥር	840
CCEETAECCAA	3.000003.000			CHICATICCCAC	IGCITAACIG	040
GCTTATCGAA	ATGTCGACTG	AGAACTTCAG	GGTGAGTTTG	GGGACCCTTG	ATTGTTCTTT	900
C	ATTGTAAAAT	TCATCTTATA	TGGAGGGGGC	$\lambda \lambda \lambda cmmmmc\lambda$	CCCTCTTCTTC	
		ICAIGITATA	TGGAGGGGC	MANGITITEM	GGGTGT.T.C.T.T.	960
TAGAATGGGA	AGATGTCCCT	TGTATCACCA	TGGACCCTCA	TGATAATTTT	C T	1020
CHUTCH & CTC	TGTTGACAAC	C 3 MMCMCMCC	mamma mmmma		CTTCTTCM	
CITICIACIC	IGIIGACAAC	CATTGTCTCC	TCTTATTTC	TTTTCATTTT	CTGTAACTTT	1080
TTCGTTAAAC	TTTAGCTTGC	ATTTGTAACG	A A ጥጥጥጥጥ A A	T	ጥጥጥ ያ ጥጥጥር ጥር	1140
3 C 3 MMCM 3 3 C	ma commones			11011011110	TIMITIGIC	T140
AGATTGTAAG	TACTTTCTCT	AATCACTTTT	TTTTCAAGGC	AATCAGGGTA	TATTATATTG	1200
TACTTCAGCA	CAGTTTTAGA	CAACAATTCT	αααππααπαα	TCATAACCTA	CAAMAMMMCM	
003.003.003		GIA CANTAGE		TOMINGIA	GAATATITCT	1260
GCATATAAAT	TCTGGCTGGC	GTGGAAATAT	TCTTATTGGT	AGAAACAACT	ACATCCTGGT	1320
$C\Delta TC\Delta TCCTC$	CCTTTCTCTT	ma mccmma ca	3 MC 3 M3 M3 C3	COCOODO	mca cca ma a a	
CHICHICCIG	CCITICICIT	IAIGGIIACA	AIGAIAIACA	CIGITIGAGA	TGAGGATAAA	1380
ATACTCTGAG	TCCAAACCGG	GCCCCTCTGC	TAACCATGTT	CATGCCTTCT	ተርተተ	1440
A C A C C TT C C TT C	CCCAACCECC	TCCTTCTTCT	CCCCCCCC T	G3.FFFFFFGGGG	1011111001	
ACAGCICCIG	GGCAACGTGC	TGGTTGT	GCTGTCTCAT	CAT"I"I"IGGCA	AGAATTGGCC	1500
GCAAGCTTCT	GCAGCATCGT	T	CTCTCTCTCA	CACACAAAACA	CTD TOTOCOCO	
						1560
GAGAATATGG	GCCAGACTGT	TACCACTCCC	TTAAGTTTGA	CCTTAGGTCA	CTGGAAAGAT	1620
GTCGAGCGGA	TCGCTCACAA	CCACTCCCTA	CATCTCAACA	ACACACCTTC	CCTTACCTTAC	
GICGIIGCGGA	ICGCICACAA	CCAGICGGIA	GAIGICAAGA	AGAGACGIIG	GGTTACCTTC	1680
TGCTCTGCAG	AATGGCCAAC	CTTTAACGTC	GGATGGCCGC	GAGACGGCAC	CTTTAACCGA	1740
CACCTCATCA	CCCAGGTTAA	CAMCAACCMC	mmmmc x ccmc	CCCCCCAMCC	202000202	
GACCICATCA	CCCAGGITAA	GATCAAGGTC	TTTTCACCTG	GCCCGCATGG	ACACCCAGAC	1800
CAGGTCCCCT	ACATCGTGAC	CTGGGAAGCC	ጥጥርGCጥጥጥጥር	ACCCCCCTCC	CTGGGTCAAG	1860
CCCMMMCMAC	7.000000	macacamaca	compounds.	CCCCCCCCC	CIOCCICANO	
CCCITIGIAC	ACCCTAAGCC	TCCGCCTCCT	CTTCCTCCAT	CCGCCCCCGTC	TCTCCCCCTT	1920
GAACCTCCTC	GTTCGACCCC	GCCTCGATCC	T	CAGCCCTCAC	T	1980
00000003330	CE3333CC		10001111110	CHOCCCICAC	ICCLICICIA	
GGCGCCAAAC	CTAAACCTCA	AGTTCTTTCT	GACAGTGGGG	GGCCGCTCAT	CGACCTACTT	2040
ACAGAAGACC	CCCCGCCTTA	TACCCACCCA	ACACCACCCC	CTTCCCACAC	CCACCCAAAM	2100 :
	CCCCCCTIA	INGGGACCCA	AGACCACCC	CITCCGACAG	GGACGGAAAT	2100 '
GGTGGAGAAG	CGACCCCTGC	GGGAGAGGCA	CCGGACCCCT	CCCCAATGGC	ATCTCGCCTA	2160
CGTGGGAGAC	GGGAGCCCCC	mamacaca x a	TO CACTA COT	CCCACCCAMM	CCCCCTCCCC	
COLOGORIGHE	GGGAGCCCCC	TGTGGCCGAC	ICCACIACCI	CGCAGGCAII	CCCCTCCCC	2220
GCAGGAGGAA	ACGGACAGCT	TCAATACTGG	CCGTTCTCCT	CTTCTGACCT	TTACAACTGG	2280
מיח ת תיח ת ת ת ת ב	ACCCTTCTTT	THE CHEST ACTOR	CCACCONAAAC	mca ca comor	Clare and and	
PUPPULITATIA	ACCCITCITI	TICIGAAGAI	CCAGGTAAAC	TGACAGCTCT	GATCGAGTCT	2340
GTTCTCATCA	CCCATCAGCC	CACCTGGGAC	GACTGTCAGC	AGCTGTTGGG	GACTCTGCTG	2400
ACCCCACAAC	77777777	COMOCHUM	C) CCCT) C))	10000000000	0110101010	
ACCEGAGAAG	AAAAACAACG	GGTGCTCTTA	GAGGCTAGAA	AGGCGGTGCG	GGGCGATGAT	2460
GGGCGCCCA	CTCAACTGCC	CAATGAAGTC	GATGCCGCTT	TTCCCCTCCA	GCGCCCAGAC	2520
mcccamma ca	667.6667.666	3.663.6663.36	61110000011	11000010011	CCCCCAGAC	
IGGGATTACA	CCACCCAGGC	AGGACGCAAC	CACCTAGTCC	ACTATCGCCA	GTTGCTCCTA	2580
GCGGGTCTCC	AAAACGCGGG	CAGAAGCCCC	ACCA ATTTCC	CCAAGGTAAA	ACCAATAACA	2640
CAACCCCCCA	1.000 CECEGO				HOOMETTE	
	ATGAGTCTCC					2700
TACACTCCTT	ATGACCCTGA	GGACCCAGGG	CDDCDDDCTD	ልጥርጥርጥርጥ አጥ	CITCHITH A TIME	2760
magaz amama			CIMIOIMMICIII	HIGIOICIAL	GICITICATI	
TGGCAGTCTG	CCCCAGACAT	'I'GGGAGAAAG	TTAGAGAGGT	TAGAAGATTT	AAAAAACAAG	2820
ACGCTTGGAG	ATTTGGTTAG	AGAGGCAGAA	ΔΑΓΑΤΌΤΤΤΑ	ATAAACGAGA	AACCCCGGAA	2880
CAAACACACAC	770007003	C3 C3 C3 C3 C3	030033333	330330000	TACCCCCCAA	2000
ODADADAGG	AACGIAICAG	GAGAGAAACA	GAGGAAAAAG	AAGAACGCCG.	TAGGACAGAG	2940
GATGAGCAGA	AAGAGAAAGA	AAGAGATCGT	AGGAGACATA	GAGAGATGAG	CAAGCTATTG	3000
CCCACMCMCC	TEN CECCA CA	23.1.2.2.2.2	10101010111		CILICCIATIO	3000
GCCACTGTCG	TTAGTGGACA	GAAACAGGA'I	AGACAGGGAG	GAGAACGAAG	GAGGTCCCAA	3060
CTCGATCGCG	ACCAGTGTGC	CTACTGCAAA	GAAAAGGGGC	ACTGGGGTAA	AGATTGTCCC	3120
7707770070	Clearence			22222222	MORITGICCC	3120
AAGAAACCAC	GAGGACCTCG	GGGACCAAGA	CCCCAGACC'I'	CCCTCCTGAC	CCTAGATGAC	3180
TAGGGAGGTC	AGGGTCAGGA	GCCCCCCCT	GAACCCAGGA	TAACCCTCAA	AGTCGGGGG	3240
Chaccomon	SSEESSES SEE	55555555		TIMICCCICIMI	DDDDDDA	3240
CAACCCGTCA	CCTTCCTGGT	AGATACTGGG	GCCCAACACT	CCGTGCTGAC	CCAAAATCCT GCGGTATCGC	3300
GGACCCCTAA	GTGATAAGTC	TECCTECETE	CAAGGGGCTA	CTCCACCAAA	CCCCTATCCC	3360
mcca coa coa	700001	100010010	2.110000011	TOUR COMM	CCGGIAICGC	3300
TGGACCACGG	ATUGCAAAGT	ACATCTAGCT	ACCGGTAAGG	TCACCCACTC	TTTCCTCCAT	3420
GTACCAGACT	GTCCCTATCC	TOTOTOTOTO	$\Delta G \Delta G \Delta T T T T C C$	TCDCTDDDCT	AAAAGCCCAA	3100
100000000	CICCIAICC	LCIGITAGGA	JULIADADA	IGACIAAACI	AMAGCCCAA	3480
ATCCACTTTG	AGGGATCAGG	AGCTCAGGTT	ATGGGACCAA	'I'GGGGCAGCC	CCTGCAAGTG	3540
ጥጥር <u>አ</u> ርርር ጥ አ	עטע עטעשעשע איי	TCACCATOCC	CHACAMCACA	CCMCXXXXCX	GCCAGATGTT	3600
LIGACCCIAA	ALALAGAAGA	TOMOCHICGG	CINCALGAGA	CCICAAAAGA	GCCAGATGTT	3600
TCTCTAGGGT	CCACATGGCT	GTCTGATTTT	CCTCAGGCCT	GGGCGGAAAC	CGGGGGCATG	3660
CCACTCCCAC	THECCENTRE	mccmcmcxmc	A M A C C M C M C A	AACCAACCEC	ma coccocc	2700
DAJOULGGCAG	CHAROCA	TCCTCTGATC	MIMCCICIGA	PURCHACCIC	TACCCCCGTG CCACATACAG	3720
TCCATAAAAC	AATACCCCAT	GTCACAAGAA	GCCAGACTGG	GGATCAAGCC	CCACATACAG	3780
ACACTCTTCC	ACCACCCA AM	A CELCCENA CCC	macca amaca	CCMCC33C3C	GCCCTGCTA	3,00
DOLLOTON	ACCAGGGAAT	ACTGGTACCC	TGCCAGICCC	CCIGGAACAC	GCCCCTGCTA	3840
CCCGTTAAGA	AACCAGGGAC	TAATGATTAT	AGGCCTGTCC	AGGATCTGAG	AGAAGTCAAC	3900
AACCCCCCCCC	770707000	CCCCACCCC	CCCAACCCCC	707700000		3,00
SOLDSON	MAGACATCCA	CCCCACCGTG	CCCAACCCT"I"	ACAACCTCTT	GAGCGGGCTC	3960
CCACCGTCCC	ACCAGTGGTA	CACTGTGCTT	GATTTAAAGG	ATGCCTTTTT	CTGCCTGAGA	4020
CTCCACCCC	CONOMOROGO	momommacc:		03030001111	CIUCUIGAGA	4020
CICCACCCCA	CCAGTCAGCC	TCTCTTCGCC	TTTGAGTGGA	GAGA'I'CCAGA	GATGGGAATC	4080
					_	-

Figure 8. hCMV+intronkaSD Sequence

тсассасаат	TCA CCTCCA C	CACACMCCCA	C) CCCMMMC)	1111010000	CACCCTGTTT	
CATCACCCAC	TGCACAGAGA	CAGACTCCCA	CAGGGTTTCA	AAAACAGTCC	CACCCTGTTT	•
CTACACTACC	TCCATCAGAGA	ACTAGCAGAC	COCACCATCC	AGCACCCAGA	CTTGATCCTG	4200
ACTCGGGGGGG	TGGATGACTT	ACTGCTGGCC	GCCACTTCTG	AGCTAGACTG	CCAACAAGGT	4260
CANATOTOCC	TGTTACAAAC	CCTAGGGAAC	CTCGGGTATC	GGGCCTCGGC	CAAGAAAGCC	4320
CHAMITICCC	AGAAACAGGT	CAAGTATCTG	GGGTATCTTC	TAAAAGAGGG	TCAGAGATGG	4380
CIGACIGAGG	CCAGAAAAGA	GACTGTGATG	GGGCAGCCTA	CTCCGAAGAC	CCCTCGACAA	4440
CTAAGGGAGT	TCCTAGGGAC	GGCAGGCTTC	TGTCGCCTCT	GGATCCCTGG	GTTTGCAGAA	4500
ATGGCAGCCC	CCTTGTACCC	TCTCACCAAA	ACGGGGACTC	TGTTTAATTG	GGGCCCAGAC	4560
CAACAAAAGG	CCTATCAAGA	AATCAAGCAA	GCTCTTCTAA	CTGCCCCAGC	CCTGGGGTTG	4620
CCAGATTTGA	CTAAGCCCTT	TGAACTCTTT	GTCGACGAGA	AGCAGGGCTA	CGCCAAAGGT	4680
GTCCTAACGC	AAAAACTGGG	ACCTTGGCGT	CGGCCGGTGG	CCTACCTGTC	CAAAAAGCTA	4740
GACCCAGTAG	CAGCTGGGTG	GCCCCCTTGC	CTACGGATGG	TAGCAGCCAT	TGCCGTACTG	4800
ACAAAGGATG	CAGGCAAGCT	AACCATGGGA	CAGCCACTAG	TCATTCTGGC	CCCCCATGCA	4860
GTAGAGGCAC	TAGTCAAACA	ACCCCCGAC	CGCTGGCTTT	CCAACGCCCG	GATGACTCAC	4920
TATCAGGCCT	TGCTTTTGGA	CACGGACCGG	GTCCAGTTCG	GACCGGTGGT	ACCCCTGAAC	4980
CCGGCTACGC	TGCTCCCACT	GCCTGAGGAA	GGGCTGCAAC	ACAACTGCCT	TGATATCCTG	5040
GCCGAAGCCC	ACGGAACCCG	ACCCGACCTA	ACGGACCAGC	CGCTCCCAGA	CGCCGACCAC	5100
ACCTGGTACA	CGGATGGAAG	CAGTCTCTTA	CAAGAGGGAC	AGCGTAAGGC	GGGAGCTGCG	5160
GTGACCACCG	AGACCGAGGT	AATCTGGGCT	AAAGCCCTGC	CAGCCGGGAC	ATCCCCTCAG	5220
CGGGCTGAAC	TGATAGCACT	CACCCAGGCC	CTAAAGATGG	CAGAAGGTAA	GAACCTAAAT	5280
GTTTATACTG	ATAGCCGTTA	TGCTTTTGCT	ACTGCCCATA	TCCATGGAGA	AATATACAGA	5340
AGGCGTGGGT	TGCTCACATC	AGAAGGCAAA	GAGATCAAAA	ATAAAGACGA	GATCTTGGCC	5400
CTACTAAAAG	CCCTCTTTCT	GCCCAAAAGA	CTTAGCATAA	TCCATTGTCC	AGGACATCAA	5460
AAGGGACACA	GCGCCGAGGC	TAGAGGCAAC	CGGATGGCTG	ACCAAGCGGC	CCGAAAGGCA	5520
GCCATCACAG	AGACTCCAGA	CACCTCTACC	CTCCTCATAG	AAAATTCATC	ACCCTACACC	5580
TCAGAACATT	TTCATTACAC	AGTGACTGAT	ATAAAGGACC	TAACCAAGTT	GGGGGGCAMM	
TATGATAAAA	CAAAGAAGTA	ТТСССТСТАС	CAAGGAAAAC	CTGTGATGCC	TCACCACTOR	5640 5700
ACTTTTGAAT	TATTAGACTT	TCTTCATCAG	CTGACTCACC	TCACCTTCTC	AAAAATCAAC	5700
GCTCTCCTAG	AGAGAAGCCA	CAGTCCCTAC	TACATGCTGA	ACCGGGATCG	A A C A CTC A A A	5760 5320
AATATCACTG	AGACCTGCAA	ACCTTCTCCA	CAACTCAACG	CCACCAACTC	MACACICAAA MCCCCCMM3 3 3	5820
CAGGGAACTA	GGGTCCGCGG	GCATCGGCCC	CCCACTCATT	GGGAGATCGA	TOCCGITAAA.	5880
ATAAAGCCCG	GATTGTATGG	СТАТАААТАТ	CTTCTACTT	TTATACATAC	COMMUNICACCO	5940
TGGATAGAAG	CCTTCCCAAC	CAAGAAAGAA	ACCCCCAACC	TCCTAACCAA	CAACCEACEA	6000
GAGGAGATCT	TCCCCAGGTT	CCCCATCCCT	CACCUATUTCC	CAACTCACAA	GAAGCTACTA TECCCCCTACTA	6060
TTCGTCTCCA	AGGTGAGTCA	CACACTGGCC	CAGGIATIGG	CCARROAMEC	CARAGECTIGCC	6120
TGTGCATACA	GACCCCAAAG	CTCACCCCAC	CENCINING	GGATTGATTG	GAAATTACAT	6180
ACTTTAACTA	AATTAACGCT	TCCAGGCCAG	GIAGAAAGAA	CCCTCCTCCT	CATCAAGGAG	6240
GCCCTGTACC	GAGCCCGCAA	CACCCCCCCC	CCCCAMCCCC	GGGTGCTCCT	ACTCCCCTTA	6300
TATEGEGECAC	CCCCGCCCT	TOTAL A A COURT	CCCCATGGCC	TCACCCCATA	TGAGATCTTA	6360
AGCCCCTCTC	TCCAAGCTCA	TGTAAACTTC	CCTGACCCTG	ACATGACAAG	AGTTACTAAC	6420
CCTCTGGCGG	CAGCCTACCA	ACAACAACTC	CICIACTIAG	TCCAGCACGA	AGTCTGGAGA	6480
GTCGGCGACA	CAGCCIMCCA	CCCCCCAACTG	GACCGACCGG	TGGTACCTCA	CCCTTACCGA	6540
CCACCTTACA	CAGTGTGGGT CAGTCCTGCT	CACCACACAC	CAGACTAAGA	ACCTAGAACC	TCGCTGGAAA	6600
TGGATACACG	CCCCCCACCT	GACCACCCCC	ACCGCCCTCA	AAGTAGACGG	CATCGCAGCT	6660
ACATEGECECE	CCGCCCACGT	GAAGGCIGCC	GACCCCGGGG	GTGGACCATC	CTCTAGACTG	6720
TAATCCCCTT	TTCAACGCTC	TCAAAACCCC	TTAAAAATAA	GGTTAACCCG	CGAGGCCCCC	6780
CGGCCCAGCC	AATTCTTCTG	ATGCTCAGAG	GGGTCAGTAC	TGCTTCGCCC	GGCTCCAGTG	6840
AAGTAGCGAC	GGCCACCATG	AAAACATTTA	ACATTTCTCA	ACAAGATCTA	GAATTAGTAG	6900
CAATTCCTAC	AGAGAAGATT	ACAATGCTTT	ATGAGGATAA	TAAACATCAT	GTGGGAGCGG	6960
CACTAACTC	GAAAACAGGA	GAAATCATTT	CGGCAGTACA	TATTGAAGCG.	TATATAGGAC	7020
VALUACIGI.	TTGTGCAGAA	GCCATTGCGA	TTGGTAGTGC	AGTTTCGAAT	GGACAAAAGG	7080
CACTCCTAACAC	GATTGTAGCT	GTTAGACACC	CTTATTCTGA	CGAAGTAGAT	AGAAGTATTC	7140
THCHCHHAN Y W	TCCTTGTGGT	ATGTGTAGGG	AGTTGATTTC	AGACTATGCA	CCAGATTGTT	7200
TIGIGITAAT	AGAAATGAAT	GGCAAGTTAG	TCAAAACTAC	GATTGAAGAA	CTCATTCCAC	7260
*CUVUIUIUC	CCGAAATTAA	AAGTTTTACC	ACCAAGC'I"I'A	TCGAATTC		7308

Figure 9. FBdelPASAF Sequence

Figure 9. FBdelPASAF Sequence

	AACCCGACAG	GACTATAAAG	ATACCAGGCG	TTTCCCCCTG	GAAGCTCCCT	4140
CGTGCGCTCT	CCTGTTCCGA	CCCTGCCGCT	TACCGGATAC	CTGTCCGCCT	TTCTCCCTTC	4200
GGGAAGCGTG	GCGCTTTCTC	AATGCTCACG	CTGTAGGTAT	CTCAGTTCGG	TGTAGGTCGT	4260
TCGCTCCAAG	CTGGGCTGTG	TGCACGAACC	CCCCGTTCAG	CCCGACCGCT	GCGCCTTATC	4320
CGGTAACTAT	CGTCTTGAGT	CCAACCCGGT	AAGACACGAC	TTATCGCCAC	TGGCAGCAGC	4380
CACTGGTAAC	AGGATTAGCA	GAGCGAGGTA	TGTAGGCGGT	GCTACAGAGT	TCTTGAAGTG	4440
GTGGCCTAAC	TACGGCTACA	CTAGAAGGAC	AGTATTTGGT	ATCTGCGCTC	TGCTGAAGCC	4500
AGTTACCTTC	GGAAAAAGAG	TTGGTAGCTC	TTGATCCGGC	AAACAAACCA	CCGCTGGTAG	4560
CGGTGGTTTT	TTTGTTTGCA	AGCAGCAGAT	TACGCGCAGA	AAAAAAGGAT	CTCAAGAAGA	4620
TCCTTTGATC	TTTTCTACGG	GGTCTGACGC	TCAGTGGAAC	GAAAACTCAC	GTTAAGGGAT	4680
TTTGGTCATG	AGATTATCAA	AAAGGATCTT	CACCTAGATC	CTTTTAAATT	AAAAATGAAG	4740
TTTTAAATCA	ATCTAAAGTA	TATATGAGTA	AACTTGGTCT	GACAGTTACC	AATGCTTAAT	4800
CAGTGAGGCA	CCTATCTCAG	CGATCTGTCT	ATTTCGTTCA	TCCATAGTTG	CCTGACTCCC	4860
CGTCGTGTAG	ATAACTACGA	TACGGGAGGG	CTTACCATCT	GGCCCCAGTG	CTGCAATGAT	4920
ACCGCGAGAC	CCACGCTCAC	CGGCTCCAGA	TTTATCAGCA	ATAAACCAGC	CAGCCGGAAG	4980
GGCCGAGCGC	AGAAGTGGTC	CTGCAACTTT	ATCCGCCTCC	ATCCAGTCTA	TTAATTGTTG	5040
CCGGGAAGCT	AGAGTAAGTA	GTTCGCCAGT	TAATAGTTTG	CGCAACGTTG	TTGCCATTGC	5100
TACAGGCATC	GTGGTGTCAC	GCTCGTCGTT	TGGTATGGCT	TCATTCAGCT	CCGGTTCCCA	5160
ACGATCAAGG	CGAGTTACAT	GATCCCCCAT	GTTGTGCAAA	AAAGCGGTTA	GCTCCTTCGG	5220
TCCTCCGATC	GTTGTCAGAA	GTAAGTTGGC	CGCAGTGTTA	TCACTCATGG	TTATGGCAGC	5280
ACTGCATAAT	TCTCTTACTG	TCATGCCATC	CGTAAGATGC	TTTTCTGTGA	CTGGTGAGTA	5340
CTCAACCAAG	TCATTCTGAG	AATAGTGTAT	GCGGCGACCG	AGTTGCTCTT	GCCCGGCGTC	5400
AATACGGGAT	AATACCGCGC	CACATAGCAG	AACTTTAAAA	GTGCTCATCA	TTGGAAAACG	5460
TTCTTCGGGG	CGAAAACTCT	CAAGGATCTT	ACCGCTGTTG	AGATCCAGTT	CGATGTAACC	5520
CACTCGTGCA	CCCAACTGAT	CTTCAGCATC	TTTTACTTTC	ACCAGCGTTT	CTGGGTGAGC	5580
AAAAACAGGA	AGGCAAAATG	CCGCAAAAAA	GGGAATAAGG	GCGACACGGA	AATGTTGAAT	5640
ACTCATACTC	TTCCTTTTTC	AATATTATTG	AAGCATTTAT	CAGGGTTATT	GTCTCATGAG	5700
CGGATACATA	TTTGAATGTA	TTTAGAAAAA	TAAACAAATA	GGGGTTCCGC	GCACATTTCC	5760
CCGAAAAGTG	CCACCTGACG	TCTAAGAAAC	CATTATTATC	ATGACATTAA	CCTATAAAAA	5820
TAGGCGTATC	ACGAGGCCCT	TTCGTCTCGC	GCGTTTCGGT	GATGACGGTG	AAAACCTCTG	5880
ACACATGCAG	CTCCCGGAGA	CGGTCACAGC	TTGTCTGTAA	GCGGATGCCG	GGAGCAGACA	5940
AGCCCGTCAG	GGCGCGTCAG	CGGGTGTTGG	CGGGTGTCGG	GGCTGGCTTA	ACTATGCGGC	6000
ATCAGAGCAG	ATTGTACTGA	GAGTGCAC				6028

Figure 10. FBdelPMOSAF Sequence

		•				
CATATGCGGT	GTGAAATACC	GCACAGATGC	GTA AGGAGA A	א אייז מרככת איי	' CAGGCGCCAT	
TCGCCATTCA	GGCTGCGCAA	CTGTTGGGAA	GGGCGATCGG	TGCGGGCCTC	TTCGCTATTA	60
CGCCAGCTGG	CGAAAGGGGG	ATGTGCTGCA	ACCCCATTAA	CTTCCCTA A C	GCCAGGGTTT	•
TCCCAGTCAC	GACGTTGTAA	AACGACGGCC	ACTICA ATTICC	CATTOOCIAC	ATTTGTTAAA	180
GACAGGATCT	CAGTAGTCCA	GGCTTTAGTC	CTGACTCAAC	JATTAGIICA	GCTAAAACCA	240
CTAGAATACG	AGCCACAATA	DOCTITAGIC	TIGACICAAC	TTTCCACAAAA	AGGGGGGAAT	300
GAAAGACCCC	ACCAAATTGC	THINCCOMON	ACCCCCACMA	1 I CCAGAAAA	GCAAGGCATG	360
GAAAAATACC	AAACCAAGAA	TAGCCIGAT	CACAMCAACC	ACGCCATTTT	GAAAACAGCT GAAAACAGCT	420
AACGTTGGGC	CAAACAGGAT	ATCTCCCCTC	A CCA CERENCO	GCGGGTACAC	GAAAACAGCT GGGGCCAAGA	480
ACAGATGGTC	ACCCCCCTTC	GGCCCCGGCC	AGCAGTTTCG	336363555	GGGGCCAAGA	540
GCCCCAACCC	TCVCCCVCTTC	CTTAAGACCC	AMCACAMO	AACAGATGGT	CCCCAGATAT	600
TCAAATCACC	CTCTCCCTTA	CTTAAGACCC	ATCAGATGTT	TCCAGGCTCC	CCCAAGGACC	660
CCCCTTCTCC	TTCCCCACCT	TTTGAATTAA	CCAATCAGCC	TGCTTCTCGC	TTCTGTTCGC	720
CTCCCATACA	CTCACCCCACCT	CTATAAAAGA	GCTCACAACC	CCTCACTCGG	CGCGCCAGTC	780
CCCACTCCTC	CTGAGICGCC	CGGGTACCCG	TGTATCCAAT	AAATCCTCTT	GCTGTTGCAT	840
CCCCCCCCCC	mcammecce.	TCCTTGGGAG	GGTCTCCTCA	GAGTGATTGA	CTACCCGTCT	900
ACCCACCACC	CCCACCTAGGG	GCTCGTCCGG	GATCTGGAGA	CCCCTGCCCA	GGGACCACCG	960
ACCUACCACC	GGGAGGTAAG	CTGGCCAAGA	TCTTATATGG	GGCACCCCG	CCCCTTGTAA	1020
ACTICCCIGA	CCCTGACATG	ACAAGAGTTA	CTAACAGCCC	CTCTCTCCAA	GCTCACTTAC	1080
AGGCTCTCTA	CTTAGTCCAG	CACGAAGTCT	GGAGACCTCT	GGCGGCAGCC	TACCAAGAAC	1140
AACTGGACCG	ACCGGTGGTA	CCTCACCCTT	ACCGAGTCGG	CGACACAGTG	TGGGTCCGCC	1200
GACACCAGAC	TAAGAACCTA	GAACCTCGCT	GGAAAGGACC	TTACACAGTC	CTGCTGACCA	1260
CCCCCACCGC	CCTCAAAGTA	GACGGCATCG	CAGCTTGGAT	ACACGCCGCC	CACGTGAAGG	1320
CTGCCGACCC	CGGGGGTGGA	CCATCCTCTA	GACTGACATG	GCGCGTTCAA	CCCTCTCAAA	1380
ACCCCTTAAA	AATAAGGTTA	ACCCGCGAGG	CCCCCTAATC	CCCTTAATTC	TTCTCATCCT	1440
CAGAGGGGTC	AGTACTGCTT	CGCCCGGCTC	CAGTCCTCAT	CAAGTCTATA	ATATCACCTC	1500
GGAGGTAACC	AATGGAGATC	GGGAGACGGT	ATGGGCAACT	TCTGGCAACC	ACCCTCTCTC	1560
GACCTGGTGG	CCTGACCTTA	CCCCAGATTT	ATGTATGTTA	GCCCACCATG	CACCATCTTA	1620
TTGGGGGCTA	GAATATCAAT	CCCCTTTTTC	TTCTCCCCCG	GGGCCCCCTT	GTTGCTC AGG	1680
GGGCAGCAGC	CCAGGCTGTT	CCAGAGACTG	CGAAGAACCT	TTAACCTCCC	TCACCCCTCC	1740
GTGCAACACT	GCCTGGAACA	GACTCAAGCT	AGACCAGACA	ACTCATAAAT	CAAATGAGGG	1800
ATTTTATGTT	TGCCCCGGGC	CCCACCGCCC	CCGAGAATCC	AAGTCATGTG	GGGGTCCAGA	1860
CTCCTTCTAC	TGTGCCTATT	GGGGCTGTGA	GACAACCGGT	AGAGCTTACT	GGAAGCCCTC	1920
CTCATCATGG	GATTTCATCA	CAGTAAACAA	CAATCTCACC	TCTGACCAGG	CTGTCCAGGT	1980
ATGCAAAGAT	AATAAGTGGT	GCAACCCCTT	AGTTATTCGG	TTTACAGACG	CCGGGAGACG	2040
GGTTACTTCC	TGGACCACAG	GACATTACTG	GGGCTTACGT	TTGTATGTCT	CCGCACAACA	2100
TCCAGGGCTT	ACATTTGGGA	TCCGACTCAG	ATACCAAAAT	CTAGGACCCC	GCGTCCCA AT	2160
AGGGCCAAAC	CCCGTTCTGG	CAGACCAACA	GCCACTCTCC	AAGCCCAAAC	ርጥርጥጥል ልርጥር	2220
GCCTTCAGTC	ACCAAACCAC	CCAGTGGGAC	TCCTCTCTCC	CCTACCCAAC	TTCCACCCC	2280
GGGAACGGAA	AATAGGCTGC	TAAACTTAGT	AGACGGAGCC	TACCAAGCCC	TCAACCTCAC	2340
CAGTCCTGAC	AAAACCCAAG	AGTGCTGGTT	GTGTCTAGTA	GCGGGACCCC	CCTACTACGA	2400
AGGGGTTGCC	GTCCTGGGTA	CCTACTCCAA	CCATACCTCT	GCTCCAGCCA	ACTGCTCCGT	2460
GGCCTCCCAA	CACAAGTTGA	CCCTGTCCGA	AGTGACCGGA	CAGGGACTCT	GCATAGGAGC	2520
AGTTCCCAAA	ACACATCAGG	CCCTATGTAA	TACCACCCAG	ACAAGCAGTC	GAGGGTCCTA	2580
TTATCTAGTT	GCCCCTACAG	GTACCATGTG	GGCTTGTAGT	ACCGGGCTTA	CTCCATGCAT	2640
CTCCACCACC	ATACTGAACC	TTACCACTGA	TTATTGTGTT	CTTGTCGAAC	TCTGGCCAAG	2700
AGTCACCTAT	CATTCCCCCA	GCTATGTTTA	CGGCCTGTTT	GAGAGATCCA	ACCGACACAA	2760
AAGAGAACCG	GIGTCGTTAA	CCCTGGCCCT	ATTATTGGGT	GGACTAACCA	TGGGGGGAAT	2920
TGCCGCTGGA	ATAGGAACAG	GGACTACTGC	TCTAATGGCC	ACTCAGCAAT	TCCAGCAGCT	2880
CCAAGCCGCA	GTACAGGATG	ATCTCAGGGA	GGTTGAAAAA	TCAATCTCTA	ACCTAGAAAA	2940
GTCTCTCACT	TCCCTGTCTG	AAGTTGTCCT	ACAGAATCGA	AGGGGCCTAG	ACTTGTTATT	3000
TCTAAAAGAA	GGAGGGCTGT	GTGCTGCTCT	AAAAGAAGAA	TGTTGCTTCT	ATGCGGACCA	3060
ACHCHURCAC	GTGAGAGACA	GCATGGCCAA	ATTGAGAGAG	AGGCTTAATC	AGAGACAGAA	3120
ACTGTTTGAG	TCAACTCAAG	GATGGTTTGA	GGGACTGTTT	AACAGATCCC	CTTGGTTTAC	3180
ACCOMMODITION	TCTACCATTA	TGGGACCCCT	CATTGTACTC	CTAATGATTT	TGCTCTTCGG	3240
ACCCTGCATT	CTTAATCGAT	TAGTTCAATT	TGTTAAAGAC	AGGATCTCAG	TAGTCCAGGC CATAGGGCGC	3300
TTTAGTCCTG	ACTCAACAAT	ACCACCAGCT	AAAGCCTATA	GAGTACGAGC	CATAGGGCGC	3360
CIAGIGITGA	CAATTAATCA	TCGGCATAGT	ATACGGCATA	GTATAATACG	ACTCACTATA	3420
GGAGGGCCAC	CATGGCCAAG	TTGACCAGTG	CCGTTCCGGT	GCTCACCGCG	CGCGACGTCG	3480
CCGGAGCGGT	CGAGTTCTGG	ACCGACCGGC	TCGGGTTCTC	CCGGGACTTC	GTGGAGGACG	3540
ACTICGCCGG	TGTGGTCCGG	GACGACGTGA	CCCTGTTCAT	CAGCGCGGTC	GTGGAGGACG CAGGACCAGG	3600
AGTGGTCGGA	GGTCGTGTCC	ACGAACTTCC	GGGACGCCTC	CGGGCCGGCC	ATGACCGAGA	3720
TUGGUGAGUA	GCCGTGGGGG	CGGGAGTTCG	CCCTGCGCGA	CCCGGCCGGC	AACTCCCTCC	3780
ACTICGIGGC	CGAGGAGCAG	GACTGANNNN	CGGACCGGTC	GACTTGTTAA	CTTGTTTATT	3840
GCAGCTTATA	ATGGTTACAA	ATAAAGCAAT	AGCATCACAA	ATTTCACAAA	TAAAGCATTT	3900
TTTTCACTGC	ATTCTAGTTG	TGGTTTGTCC	AAACTCATCA	ATGTATCTTA	TCATGTCTGG	3960
ATCCAGATCT	GGGCCCATGC	GGCCGCGGAT	CGATNNNNAC	ATGTGAGCAA	AAGGCCACCA	1020
AAAGGCCAGG	AACCGTAAAA	AGGCCGCGTT	GCTGGCGTTT	TTCCATAGGC	TCCGCCCCCC	4080

Figure 10. FBdelPMOSAF Sequence

TGACGAGCAT	CACAAAAATC	GACGCTCAAG	TCAGAGGTGG	CCANACCCCA	CAGGACTATA		
AAGATACCAG	GCGTTTCCCC	CTGGAAGCTC	CCTCGTGCGC	TCTCCTGTTC		-	4140
GCTTACCGGA	TACCTGTCCG	CCTTTCTCCC	TTCGGGAAGC	GTGGCGCTTT	CGACCCTGCC		4200
ACGCTGTAGG	TATCTCAGTT	CGGTGTAGGT	CGTTCGCTCC	AAGCTGGGCT	CTCAATGCTC		4260
ACCCCCCGTT	CAGCCCGACC	GCTGCGCCTT	ATCCGGTAAC	TATCGTCTTG	GTGTGCACGA		4320
GGTAAGACAC	GACTTATCGC	CACTGGCAGC	AGCCACTGGT	AACAGGATTA	AGTCCAACCC		4380
GTATGTAGGC	GGTGCTACAG	AGTTCTTGAA	GTGGTGGCCT	AACTACGGCT	-011011000110		4440
GACAGTATTT	GGTATCTGCG	CTCTGCTGAA	GCCAGTTACC	TTCGGAAAAA	ACACTAGAAG		4500
CTCTTGATCC	GGCAAACAAA	CCACCGCTGG	TAGCGGTGGT	TTTTTTGTTT	GAGTTGGTAG		4560
GATTACGCGC	AGAAAAAAG	GATCTCAAGA	AGATCCTTTG	ATCTTTTCTA	GCAAGCAGCA		4620
CGCTCAGTGG	AACGAAAACT	CACGTTAAGG	GATTTTGGTC	ATGAGATTAT	CGGGGTCTGA		4680
CTTCACCTAG	ATCCTTTTAA	ATTAAAAATG	AAGTTTTAAA		CAAAAAGGAT		4740
GTAAACTTGG	TCTGACAGTT	ACCAATGCTT	AATCAGTGAG	TCAATCTAAA GCACCTATCT	GTATATATGA		4800
TCTATTTCGT	TCATCCATAG	TTGCCTGACT	CCCCGTCGTG		CAGCGATCTG		4860
GGGCTTACCA	TCTGGCCCCA	GTGCTGCAAT	GATACCGCGA	TAGATAACTA GACCCACGCT	CGATACGGGA		4920
AGATTTATCA	GCAATAAACC	AGCCAGCCGG	AAGGGCCGAG	CGCAGAAGTG	CACCGGCTCC		4980
TTTATCCGCC	TCCATCCAGT	CTATTAATTG	TTGCCGGGAA	GCTAGAGTAA	GTCCTGCAAC		5040
AGTTAATAGT	TTGCGCAACG	TTGTTGCCAT	TGCTACAGGC	ATCGTGGTGT	GTAGTTCGCC		5100
GTTTGGTATG	GCTTCATTCA	GCTCCGGTTC	CCAACGATCA		CACGCTCGTC		5160
CATGTTGTGC	AAAAAAGCGG	TTAGCTCCTT	CGGTCCTCCG	ATCGTTGTCA	CATGATCCCC		5220
GGCCGCAGTG	TTATCACTCA	TGGTTATGGC	AGCACTGCAT	AATTCTCTTA	GAAGTAAGTT		5280
ATCCGTAAGA	TGCTTTTCTG	TGACTGGTGA	GTACTCAACC		CTGTCATGCC		5340
TATGCGGCGA	CCGAGTTGCT	CTTGCCCGGC	GTCAATACGG	AAGTCATTCT	GAGAATAGTG		5400
CAGAACTTTA	AAAGTGCTCA	TCATTGGAAA	ACGTTCTTCG	GATAATACCG	CGCCACATAG		5460
CTTACCGCTG	TTGAGATCCA	GTTCGATGTA	ACCCACTCGT	GGGCGAAAAC	TCTCAAGGAT		5520
ATCTTTTACT	TTCACCAGCG	TTTCTGGGTG	AGCAAAAACA	GCACCCAACT	GATCTTCAGC		5580
AAAGGGAATA	AGGGCGACAC	GGAAATGTTG	AATACTCATA	GGAAGGCAAA CTCTTCCTTT	ATGCCGCAAA		5640
TTGAAGCATT	TATCAGGGTT	ATTGTCTCAT	GAGCGGATAC	ATATTTGAAT	TTCAATATTA		5700
AAATAAACAA	ATAGGGGTTC	CGCGCACATT	TCCCCGAAAA	GTGCCACCTG	GTATTTAGAA		5760
AACCATTATT	ATCATGACAT	TAACCTATAA	AAATAGGCGT	ATCACGAGGC	ACGTCTAAGA		5820
CGCGCGTTTC	GGTGATGACG	GTGAAAACCT	CTGACACATG	CAGCTCCCGG	CCTTTCGTCT		5880
AGCTTGTCTG	TAAGCGGATG		ACAAGCCCGT	CAGGGCGCGT	AGACGGTCAC CAGCGGGTGT		5940
TGGCGGGTGT	CGGGGCTGGC		GGCATCAGAG	CAGGGCGCGT	TGAGAGTGCA		6000
C			OCCILICACAG	CAGALIGIAC	1 GAGAGTGCA		6060
							6061

Figure 11. FBdelPGASAF Sequence

CATATGCGGT	GTGAAATACC	GCACAGATGO	GTAAGGAGAA	AATACCGCAT	CAGGCGCCAT	
TCGCCATTCA	. GGCTGCGCAA	. CTGTTGGGAA	L GGGCGATCGG	TECCCCCCCTC	The second secon	_ 60
CGCCMGCTGG	CGAAAGGGGG	: ATGTGCTGCA	L AGGCGATTAA	GTTGGGTA AC	CCCACCCMMM	120
TCCCAGTCAC	GACGTTGTAA	. AACGACGGCC	: AGTGAATTCC	GATTAGTTCA	$\Delta \Phi \Phi$	180
GACAGGATCT	CAGTAGTCCA	. GGCTTTAGTC	: CTGACTCAAC	AATACCACCA	CCTAAAAAAA	240
CTAGAATACG	AGCCACAATA	AATAAAAGAT		TTTCCACAAAA	AGGGGGGAAT	300
GAAAGACCCC	ACCAAATTGO	TTAGCCTGAT	AGCCGCAGTA	ACCCC A TOTOL	GCAAGGCATG	360
GAAAAATACC	AAACCAAGAA	TAGAGAAGTT	CAGATCAACC	CCCCCMACAC	GAAAACAGCT	420
AACGTTGGGC	CAAACAGGAT	ATCTGCGGTG	ACCACTTTCC	CCCCCCCCCC	GGGGCCAAGA	480
ACAGATGGTC	ACCGCGGTTC	GGCCCCGGCC	CCCCCCAAC	A A C A C A MCCM	GGGGCCAAGA	540
GGCCCAACCC	TCAGCAGTTT	CTTAACACCC	ATCACATION	MACAGATGGT	CCCCAGATAT	600
TGAAATGACC	CTGTGCCTTA		CCAAMCACCC	TCCAGGCTCC	CCCAAGGACC	660
GCGCTTCTGC	TTCCCGACCT	CTATATIAM	CCAATCAGCC	TGCTTCTCGC	TTCTGTTCGC	720
CTCCGATAGA	TTCCCGAGCT CTGAGTCGCC	CCCCTTACCCC	GCTCACAACC	CCTCACTCGG	CGCGCCAGTC	780
CCGACTCGTG	GTCTCCCTCT	TCCTTCCC3	TGTATCCAAT	AAATCCTCTT	GCTGTTGCAT	840
CGGGGGGTCTT	GTCTCGCTGT	CCTCCTCCC	GGTCTCCTCA	GAGTGATTGA	CTACCCGTCT	900
ACCCACCACC	TCATTTGGGG	CECCCCAACA	GATCTGGAGA	CCCCTGCCCA	GGGACCACCG	960
CGGCCTTTGT	GGGAGGTAAG	A CTICA CCCA C	TCCCTAAGGT	ACTCGGGTCA	GACAATGGCC	1020
TACATTGTGC	TGCTCAGGTA	CACACCECAC	TGGCCACTCA	ACTGGGGATA	AATTGGAAGT	1080
AAGAGACCTT	GTATAGACCC	CCCMMACAGA	GTCAGGTAGA	AAGAATGAAC	AGAACAATTA	1140
CCTTAGCGCT	GACCAAATTA	ACCA AMAGA	CCGGTGGAAA	AGACTGGGTG	ACCCTCCTTC	1200
TTCTCTATCG	GCTTAGGGCC	CCCAMACUC	2.CTGGCCGGTT	TGGTTTAACT	CCTTATGAAA	1260
GATTTCTCCC	AGGACCACCC	CCCATACTTG	AGTCTGGAGA	AACTTTGGGT	CCCGATGATA	1320
GGGACCAGAT	TGTCTTATTT	ACTCACTTAA	AGGCTTTTAGA	AATTGTAAGG	ACCCAAATCT	1380
TCGCGCATCA	CAAAGAGGTG	TATAAGCCTG	GTACCGTAAC	AATCCCTCAC	CCGTTCCAGG	1440
GCCCATACCT	AGTGCTTGTC	AGACGCCATC	GACCCAGCAG	CCTTGAGCCT	CGGTGGAAAG	1500
GCCCATACCT	GGTGTTGCTG	ACTACCCCGA	CCGCGGTAAA	AGTCGATGGT	ATTGCTGCCT	1560
TCCAAAACAC	TTCTCACCTC	AAACCTGCAC	CACCTTCGGC	ACCAGATGAG	TCCTGGGAGC	1620
A A DA A CA A CC	TGATCATCCT	CTTAAGCTGC	GTATTCGGCG	GCGGCGGGAC	GAGTCTGCAA	1680
GULLERGE	CCCACCAGCC	CATGACCCTC	ACTTGGCAGG	TACTGTCCCA	AACTGGAGAC	1740
GIIGICIGGG	ATACAAAGGC	AGTCCAGCCC	CCTTGGACTT	GGTGGCCCAC	ACTTAAACCT	1800
TCCTCTAAAC	CCTTGGCGGC	TAGTCTTGAG	TCCTGGGATA	TCCCGGGAAC	CGATGTCTCG	1860
TEGEGAGEE	GAGTCAGACC	TCCGGACTCA	GACTATACTG	CCGCTTATAA	GCAAATCACC	1920
TACCTATCTC	TAGGGTGCAG	CTACCCTCGG	GCTAGGACTA	GAATGGCAAG	CTCTACCTTC	1980
TCCCTATACT	CCCGGGATGG	CCGGACCCTT	TCAGAAGCTA	GAAGGTGCGG	GGGGCTAGAA	2040
TCCTCAAAAG	GTAAAGAATG	GGATTGTGAG	ACCACGGGGA	CCGGTTATTG	GCTATCTAAA	2100
CAACAGTGTC	ACCTCATAAC	TGTAAAATGG	GACCAAAATA	GCGAATGGAC	TCAAAAATTT	2160
AAATTATCCA	ACCAGACCGG	CIGGIGIAAC	CCCCTTAAAA	TAGATTTCAC	AGACAAAGGA	2220
CATCCAGGGG	AGGACTGGAT	AACGGGAAAA	ACCTGGGGAT	TAAGATTCTA	TGTGTCTGGA	2280
GGTCCTGACC	TACAGTTCAC	CATTCGCTTA	AAAATCACCA	ACATGCCAGC	TGTGGCAGTA	2340
CCTCTTCCCC	TCGTCCTTGT	GGAACAAGGA	CCTCCTAGAA	CGTCCCTCGC	TCTCCCACCT	2400
GCGACTAGTG	CAAGGGAAGC	GCCACCGCCA	TCTCTCCCCG	ACTCTAACTC	CACAGCCCTG	2460
CCCACCACAG	CACAAACTCC	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	AAAACAATTG	TTACCCTAAA	CACTCCGCCT	2520
ACCAACCCAG	GCGACAGACT	CTCTTCCCCCC	GTGCAGGGG	CCTTCCTAAC	CTTAAATGCT	2580
GAAGCAATAG	GGGCCACTGA CCTCATCAGG	ACACCTICCTCC	CTTTGTTTGG	CCATGGGCCC	CCCTTATTAT	2640
GGGACCCAAG	GAAAGCTCAC	CCTCACTCAC	CECECACCA	ACCTTGACCG	GTGCCGCTGG	2700
GTGCCCTTTA	CCCATCAGCA	TCTCTCCAAA	GICICAGGAC	ACGGGTTGTG	CATAGGAAAG	2760
CATCAGTATC	TGCTCCCCTC	CAACCATACC	TCCTCCCCTAT	CCATCAATTC	CTCCGGAGAC	2820
TGCCTCTCCA	CCTCAGTTTT	TAATCACACT	1GG1GGGC1T	GCAGCACTGG	CCTCACCCCT	2880
CCTCGCATCT	ATTACTATCC	TGAAGAGT	TTCTTACACC	CCMARCAGGT	CCAGCTGATT	2940
AGGACTAAAA	GAGAGGCTGT	CTCACTTACC	CTACCTCTTT	CCTATGACAA	TTCTCACCCC	3000
GCGGGAATAG	GTACTGGTTC	AACTGCCTTA	ATTANCENC	CMAMACACOM	GGGAATCACG	3060
CTGACAAGCC	TCCAGATCGC	CATAGATGCT	CACCTCCCCC	CCCTCCAACCT	CCAGCAAGGC	3120
AAGTTAGAGG	ACTCACTGAC	TTCCCTCTCC	GACCICCGGG	TCCAAGA	CTCAGTCAGC	3180
GACTTGCTGT	TTCTAAAAGA	AGGTGGCCTC	TOTOCCCCCC	TOCAMANTAG	GAGAGGCCTT	3240
TACATAGACC	ACTCAGGTGC	AGTACGGGAC	TGTGCGGCCC	AAAAGGAAGA	GTGCTGTTT	
AAAAGACAGT	TAGAGCGCCA	GAAAAGCCAA	A A CTCCTTA TC	AACTCAAAGA	AAAACTGGAT	3360
CCTTGGTTCA	CTACCCTGCT	ATCAACCATC	CCTCCCCCCC	MAGGATGGTT	CAATAACTCC	3420
CTCATCCTCG	GGCCATGCAT	CATCAATCGA	THE REPORT OF A THE	THITACICCI	CCTTCTGTTG	3480
GTAGTCCAGG	CTTTAGTCCT	GACTCAACAA	TACCACCACC	TAAACCCONA	ACACHROCA	3540
CCATAGGGCG	CCTAGTGTTG	ACAATTAATC	ATCCCCATAC	TANAGCCIAT	AGAGTACGAG	3600
GACTCACTAT	AGGAGGCCA	CCATGGCCAA	GTTGACCACT	TATACGGCAT	AGTATAATAC	3660
GCGCGACGTC	GCCGGAGCGG	TCGAGTTCTG	GACCGACCGC	CACCCT LCCCC	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	3720
CGTGGAGGAC	GACTTCGCCG	GTGTGGTCCG	GGACGACGTG	ACCCTCTTCA	TCACCCCCCC	3780
CCAGGACCAG	GIGGIGCCGG	ACAACACCCT	GGCCTGGGTG	TEGETTECECE	CCCTCCACCA	3840
GCIGIACGCC	GAGTGGTCGG	AGGTCGTGTC	CACGAACTTC	CGGGACGCCT	CCCCCCCCCC	3900
CAIGACCGAG	ATCGGCGAGC	AGCCGTGGGG	GCGGGAGTTC	GCCCTGCGCG	ACCCCCCCCCC	3960
CAACTGCGTG	CACTTCGTGG	CCGAGGAGCA	GGACTGANNIN	NCCC racaca	CCA CEMCETTS	4020
			- OTTO T OT TEATAIN	unccod1	CGACTTGTTA	4080

Figure 11. FBdelPGASAF Sequence

ACTTGTTTAT TGCAGCTTAT AATGGTTACA AATAAAGCAA TAGCATCACA AATTTCACAA ATAAAGCATT TTTTTCACTG CATTCTAGTT GTGGTTTGTC CAAACTCATC AATGTATCTT ATCATGTCTG GATCCAGATC TGGGCCCATG CGGCCGCGGA TCGATNNNNA CATGTGAGCA AAAGGCCAGC AAAAGGCCAG GAACCGTAAA AAGGCCGCGT TGCTGGCGTT TTTCCATAGG CTCCGCCCCC CTGACGAGCA TCACAAAAAT CGACGCTCAA GTCAGAGGTG GCGAAACCCG CTCCGCCCC CTGACGAGCA TCACAAAAAT CGACGCTCAA GTCAGAGGTG GCGAAACCCG
ACAGGACTAT AAAGATACCA GGCGTTTCCC CCTGGAAGCT CCCTCGTGCG CTCTCCTGTT
CCGACCCTGC CGCTTACCGG ATACCTGTCC GCCTTTCTCC CTTCGGGAAG CGTGGCGCTT
TCTCAATGCT CACGCTGTAG GTATCTCAGT TCGGTGTAGG TCGTTCGCTC CAAGCTGGGC
TGTGTGCACG AACCCCCGT TCAGCCCGAC CGCTGCGCCT TATCCGGTAA CTATCGTCTT
GAGTCCAACC CGGTAAGACA CGACTTATCG CCACTGGCAG CAGCCACTGG TAACAGGATT
AGCAGAGCGA GGTATGTAGG CGGTGCTACA GAGTTCTTGA AGTGGTGGC TAACTACGGC 4500 4740 TACACTAGAA GGACAGTATT TGGTATCTGC GCTCTGCTGA AGCCAGTTAC CTTCGGAAAA AGAGTTGGTA GCTCTTGATC CGGCAAACAA ACCACCGCTG GTAGCGGTGG TTTTTTTGTT 4860 TGCAAGCAGC AGATTACGCG CAGAAAAAAA GGATCTCAAG AAGATCCTTT GATCTTTCT ACGGGGGTCTG ACGCTCAGTG GAACGAAAAC TCACGTTAAG GGATTTTGGT CATGAGATTA 4920 TCAAAAAGGA TCTTCACCTA GATCCTTTTA AATTAAAAAT GAAGTTTTAA ATCAATCTAA 5040 AGTATATATG AGTAAACTTG GTCTGACAGT TACCAATGCT TAATCAGTGA GGCACCTATC 5100 TCAGCGATCT GTCTATTTCG TTCATCATA GTTGCCTGAC TCCCCGTCGT GTAGATAACT ACGATACGGG AGGCCTTACC ATCTGGCCC AGTGCTGCAA TGATACCGC AGACCCACGC TCACCGGCTC CAGATTTATC AGCAATAAAC CAGCCAGCCG GAAGGGCCGA GCGCAGAAGT GTTCTGCAA CTTTATCGCC CTCCATCCAG TCTATTAATT GTTGCCGGGA AGCTAGAGTA 5160 5220 AGTAGTTCGC CAGTTAATAG TTTGCGCAAC GTTGTTGCCA TTGCTACAGG CATCGTGGTG TCACGCTCGT CGTTTGGTAT GGCTTCATTC AGCTCCGGTT CCCAACGATC AAGGCGAGTT ACATGATCCC CCATGTTGTG CAAAAAAGCG GTTAGCTCCT TCGGTCCTCC GATCGTTGTC AGAAGTAAGT TGGCCGCAGT GTTATCACTC ATGGTTATGG CAGCACTGCA TAATTCTCTT 5460 5520 ACTGTCATGC CATCCGTAAG ATGCTTTTCT GTGACTGGTG AGTACTCAAC CAAGTCATTC TGAGAATAGT GTATGCGGCG ACCGAGTTGC TCTTGCCCGG CGTCAATACG GGATAATACC GCGCCACATA GCAGAACTTT AAAAGTGCTC ATCATTGGAA AACGTTCTTC GGGGCGAAAA 5700 5760 CTCTCAAGGA TCTTACCGCT GTTGAGATCC AGTTCGATGT AACCCACTCG TGCACCCAAC TGATCTTCAG CATCTTTAC TTTCACCAGC GTTTCTGGGT GAGCAAAAAC AGGAAGGCAA AATGCCGCAA AAAAGGGAAT AAGGGCGACA CGGAAATGTT GAATACTCAT ACTCTTCCTT 5820 5880 TTTCAATATT ATTGAAGCAT TTATCAGGGT TATTGTCTCA TGAGCGGATA CATATTTGAA TGTATTTAGA AAAATAAACA AATAGGGGTT CCGCGCACAT TTCCCCGAAA AGTGCCACCT GACGTCTAAG AAACCATTAT TATCATGACA TTAACCTATA AAAATAGGCG TATCACGAGG CCCTTTCGTC TCGCGCGTTT CGGTGATGAC GGTGAAAACC TCTGACACAT GCAGCTCCCG GAGACGGTCA CAGCTTGTCT GTAAGCGGAT GCCGGGAGCA GACAAGCCCG TCAGGGCGCG TCAGCGGGTG TTGGCGGGTG TCGGGGCTGG CTTAACTATG CGGCATCAGA GCAGATTGTA CTGAGAGTGC AC 6300 6312

Figure 12. FBdelPRDSAF Sequence 1

CATATIGGGT GIGAAATACC GACAGATICO GIAAGGAGA AATACCGCAT CAGGGCCTE TCGCTATTA 1,000 CGCCACCTGG GACGGCTG ACTGTTGGGA GGGGGATTG GCGGGCCT TCGCTATTA 1,100 CGCCACCTGG GACGGTTTA AACAGGGGG ATTGTCCGA GGGGGTTTA GTTGGGTAA GGCGGGCTC TCGCTATTA 1,100 CGCCACCTGC GACGGTTTA AACAGGGGG ATTGTTGTCA AGCGGGGCT TTGCTCATATA 1,100 CGCCACCTG GACGGTTTA AACAGGGGG ATTGTTGTCA AACAGGGGGAT TCCCACATCA AATACAGGA ATTGTTAAA ACAGGGGGAT TCCCACATCA AATACAGGA ATTGCTCAA AACAGGAGA ATTGCTCAACAG ACAGGATTCA ATTTGTTAAA 240 CGGGGGAAAACACA AATACAGGA TTACCTGATA ACACGGGATTA ACACGGGATTA ACACGGGATTA ACACGGGATTA ACACGGGATTA ACACGGGATTA ACACGGGATTA ACACGGATTA CACAGATTA ACACGGATTA CACAGATTA ACACGGATTA CACAGATTA ACACGGATTA CACAGATTA ACACGGATTA CACAGATTA CACAGATTA ACACGGATTA CACAGATTA CACAGATTA CACAGATTA ACACGGATTA CACAGATTA CACAGATTA ACACGGATTA CACAGATTA ACACAGATTA ACACACAGATTA ACACAGATTA ACACACAGATTA ACACAGATTA ACACAGATTA ACACAGATTA ACACACCAGATTA ACACACAGATTA ACACACCAGATTA ACACACAGATTA ACACACCAGATTA ACACACCAGATTA ACACACCAGATTA ACACACCAGATTA ACACACCAGATTA ACACACCAGATTA ACACACCAGATTA CATATGCGGT GTGAAATACC GCACAGATGC GTAAGGAGAA AATACCGCAT CAGGCGCCAT TCGCCATTCA GGCTGCGCAA CTGTTGGGAA GGGCGATCGG TGCGGGCCTC TTCGCTATTA

Figure 12. FBdelPRDSAF Sequence

AGTTCGGTGT	AGGTCGTTCG	CTCCAAGCTG	GGCTGTGTGC	ACGAACCCCC	CGTTCAGCCC	4140
GACCGCTGCG	CCTTATCCGG	TAACTATCGT	CTTGAGTCCA	ACCCGGTAAG	ACACGACTTA	4200
TCGCCACTGG	CAGCAGCCAC	TGGTAACAGG	ATTAGCAGAG	CGAGGTATGT	AGGCGGTGCT	4260
ACAGAGTTCT	TGAAGTGGTG	GCCTAACTAC	GGCTACACTA	GAAGGACAGT	ATTTGGTATC	4320
TGCGCTCTGC	TGAAGCCAGT	TACCTTCGGA	AAAAGAGTTG	GTAGCTCTTG	ATCCGGCAAA	4380
CAAACCACCG	CTGGTAGCGG	TGGTTTTTTT	GTTTGCAAGC	AGCAGATTAC	GCGCAGAAAA	4440
AAAGGATCTC	AAGAAGATCC	TTTGATCTTT	TCTACGGGGT	CTGACGCTCA	GTGGAACGAA	4500
AACTCACGTT	AAGGGATTTT	GGTCATGAGA	TTATCAAAAA	GGATCTTCAC	CTAGATCCTT	4560
TTAAATTAAA	AATGAAGTTT	TAAATCAATC	TAAAGTATAT	ATGAGTAAAC	TTGGTCTGAC	4620
AGTTACCAAT	GCTTAATCAG	TGAGGCACCT	ATCTCAGCGA	TCTGTCTATT	TCGTTCATCC	4680
ATAGTTGCCT	GACTCCCCGT	CGTGTAGATA	ACTACGATAC	GGGAGGGCTT	ACCATCTGGC	4740
CCCAGTGCTG	CAATGATACC	GCGAGACCCA	CGCTCACCGG	CTCCAGATTT	ATCAGCAATA	4800
AACCAGCCAG	CCGGAAGGGC	CGAGCGCAGA	AGTGGTCCTG	CAACTTTATC	CGCCTCCATC	4860
CAGTCTATTA	ATTGTTGCCG	GGAAGCTAGA	GTAAGTAGTT	CGCCAGTTAA	TAGTTTGCGC	4920
AACGTTGTTG	CCATTGCTAC	AGGCATCGTG	GTGTCACGCT	CGTCGTTTGG	TATGGCTTCA	4980
TTCAGCTCCG	GTTCCCAACG	ATCAAGGCGA	GTTACATGAT	CCCCCATGTT	GTGCAAAAA	5040
GCGGTTAGCT	CCTTCGGTCC	TCCGATCGTT	GTCAGAAGTA	AGTTGGCCGC	AGTGTTATCA	5100
CTCATGGTTA	TGGCAGCACT	GCATAATTCT	CTTACTGTCA	TGCCATCCGT	AAGATGCTTT	5160
TCTGTGACTG	GTGAGTACTC	AACCAAGTCA	TTCTGAGAAT	AGTGTATGCG	GCGACCGAGT	5220
TGCTCTTGCC	CGGCGTCAAT	ACGGGATAAT	ACCGCGCCAC	ATAGCAGAAC	TTTAAAAGTG	5280
CTCATCATTG	GAAAACGTTC	TTCGGGGCGA	AAACTCTCAA	GGATCTTACC	GCTGTTGAGA	5340
TCCAGTTCGA	TGTAACCCAC	TCGTGCACCC	AACTGATCTT	CAGCATCTTT	TACTTTCACC	5400
AGCGTTTCTG	GGTGAGCAAA	AACAGGAAGG	CAAAATGCCG	CAAAAAAGGG	AATAAGGGCG	5460
ACACGGAAAT	GTTGAATACT	CATACTCTTC	CTTTTTCAAT	ATTATTGAAG	CATTTATCAG	5520
GGTTATTGTC	TCATGAGCGG	ATACATATTT	GAATGTATTT	AGAAAAATAA	ACAAATAGGG	5580
GTTCCGCGCA	CATTTCCCCG	AAAAGTGCCA	CCTGACGTCT	AAGAAACCAT	TATTATCATG	5640
ACATTAACCT	ATAAAAATAG	GCGTATCACG	AGGCCCTTTC	GTCTCGCGCG	TTTCGGTGAT	5700
GACGGTGAAA	ACCTCTGACA	CATGCAGCTC	CCGGAGACGG	TCACAGCTTG	TCTGTAAGCG	5760
	GCAGACAAGC	CCGTCAGGGC	GCGTCAGCGG		GTGTCGGGGC	5820
IGGCITAACT	ATGCGGCATC	AGAGCAGATT	GTACTGAGAG	TGCAC		5865

Figure 13. hCMV10A1 Sequence

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CLASSIFICATION OF SUBJECT MATTER C 6 C12N15/86 C12N5/10 C12N15/67 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. JOURNAL OF VIROLOGY 69 (7). 1995. Α 1 - 294086-4094. ISSN: 0022-538X. July 1995, XP002023654 LUUKKONEN B G M ET AL: "Efficiency of reinitiation of translation on human immunodeficiency virus type 1 mRNAs is determined by the length of the upstream open reading frame and by intercistronic distance." see the whole document VIROLOGY (1995), 208(1), 215-25 CODEN: Α 1-29 VIRLAX; ISSN: 0042-6822, 1 April 1995, XP002023655 HERZOG, ETIENNE ET AL: "Translation of the second gene of peanut clump virus RNA 2 occurs by leaky scanning in vitro" see the whole document Further documents are listed in the continuation of box C. Х Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docudocument referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 1 2, 02, 97 23 January 1997 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Hornig, H Fax: (+31-70) 340-3016

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